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Sir:

Transmitted herewith for filing is the patent application of:

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FOR: NOVEL STREPTOCOCCUS ANTIGENS

Enclosed are:	
\boxtimes	76 pages of specification, claims, abstract.
	Declaration and Power of Attorney.
\boxtimes	Priority Claimed.
	Certified copy of
\boxtimes	33 sheets of formal drawing.
	An assignment of the invention to
	and the assignment recordation fee.
	An associate power of attorney.
	Information Disclosure Statement, Form PTO-1449 and reference.
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Respectfully submitted,

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NOVEL STREPTOCOCCUS ANTIGENS

This application claims priority from US patent application 60/113,800 filed december 23 1998 which is herein incorporated by reference.

5 FIELD OF THE INVENTION

The present invention is related to antigens, more particularly protein antigens of streptococcus pneumoniaepathogen which are useful as vaccine components for therapy and/or prophylaxis.

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BACKGROUND OF THE INVENTION

media in infants and young children.

<u>S. pneumoniae</u> is an important agent of disease in man especially among infants, the elderly and immunocompromised persons. It is a bacterium frequently isolated from patients with invasive diseases such as bacteraemia/septicaemia, pneumonia, meningitis with high morbidity and mortality throughout the world. Even with appropriate antibiotic therapy, pneumococcal infections still result in many deaths. Although the advent of antimicrobial drugs has reduced the overall mortality from pneumococcal disease, the presence of resistant pneumococcal organisms has become a major problem in the world today. Effective pneumococcal vaccines could have a major impact on the morbidity and mortality associated with <u>S. pneumoniae</u> disease. Such vaccines would also potentially be useful to prevent otitis

Efforts to develop a pneumococcal vaccine have generally concentrated on generating immune responses to the pneumococcal capsular polysaccharide. More than 80 pneumococcal capsular serotypes have been identified on the basis of antigenic differences. The currently available pneumococcal vaccine, comprising 23 capsular polysaccharides that most frequently

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caused disease, has significant shortcomings related primarily to the poor immunogenicity of some capsular polysaccharides, the diversity of the serotypes and the differences in the distribution of serotypes over time, geographic areas and age groups. In particular, the failure of existing vaccines and capsular conjugate vaccines currently in development to protect young children against all serotypes spurres evaluation of other S. pneumoniae components. Although immunogenicity of capsular polysaccharides can be improved, serotype specificity will still represent a major limitation of polysaccharide-based vaccines. The use of a antigenically conserved immunogenic pneumococcal protein antigen, either by itself or in combination with additional components, offers the possibility of a protein-based pneumococcal vaccine.

PCT Publication number WO98/18930 published may 7 1998 entitled "Streptococcus Pneumoniae antigens and vaccines" describes certain polypeptides which are claimed to be antigenic. However, no biological activity of these polypeptides is reported.

Therefore their remains an unmet need for Streptococcus antigens that may be used as vaccine components for the prophylaxis and/or therapy of Streptococcus infection.

SUMMARY OF THE INVENTION

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

In other aspects, there are provided vectors comprising polynucleotides of the invention operably linked to an expression control region, as well as host cells transfected with said vectors and methods of producing polypeptides comprising culturing said host cells under conditions suitable for expression.

In yet another aspect, there are provided novel polypeptides encoded by polynucleotides of the invention.

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BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is the DNA sequence of BVH-3 gene; SEQ ID NO: 1.

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Figure 2 is the amino acid sequence of BVH-3 protein; SEQ ID NO: 2.

Figure 3 is the DNA sequence of BVH-11 gene; SEQ ID NO: 3.

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Figure 4 is the amino acid sequence of BVH-11 protein; SEQ ID NO: 4.

Figure 5 is the DNA sequence of BVH-28 gene; SEQ ID NO: 5.

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Figure 6 is the amino acid sequence of BVH-28 protein; **SEQ ID** NO: 6.

Figure 7 is the DNA sequence of BVH-3A gene which corresponds to the 5' terminal end of BVH-3; **SEQ ID NO: 7.**

Figure 8 is the amino acid sequence of BVH-3A protein; SEQ ID NO: 8.

Figure 9 is the DNA sequence of BVH-3B gene which corresponds to the 3' terminal end of BVH-3; **SEQ ID NO: 9.**

Figure 10 is the amino acid sequence of BVH-3B protein; SEQ ID NO: 10.

Figure 11 depicts the comparison of the predicted amino acid sequences of the BVH-3 open reading frames from WU2, RX1, JNR.7/87, SP64, P4241 and A66 S. pneumoniae strains by using the program Clustal W from MacVector sequence analysis software (version 6.5). Underneath the alignment, there is a consensus line where * and . characters indicate identical and similar amino acid residues, respectively.

Figure 12 depicts the comparison of the predicted amino acid sequences of the BVH-11 open reading frames from WU2, Rx1, JNR.7/87, SP64, P4241, A66 and SP63 S. pneumoniae strains by using the program Clustal W from MacVector sequence analysis software (version 6.5). Underneath the alignment, there is a consensus line where * and . characters indicate identical and similar amino acid residues, respectively.

Figure 13 depicts the comparison of the predicted amino acid sequences of the BVH-11 proteins from various <u>S. pneumoniae</u> strains. The degrees of identity (I) and similarity (S) were determined by using the program Clustal W from MacVector sequence analysis software (version 6.5).

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Figure 14 is a DNA sequence containing the complete BVH-3 gene (open reading frame "ORF" at nucleotides 1777 to 4896); **SEQ ID NO: 11.**

5 Figure 15 is a DNA sequence containing the complete BVH-11 gene (ORF at nucleotides 45 to 2567); SEQ ID NO: 12.

Figure 16 is a DNA sequence containing the complete BVH-11-2 gene (ORF at nucleotides 114 to 2630); SEQ ID NO: 13.

Figure 17 is the amino acid sequence of BVH-11-2 protein; SEQ ID NO: 14.

Figure 18 is the DNA sequence of SP63 BVH-3 gene; SEQ ID NO:15.

Figure 19 is the amino acid sequence of SP63 BVH-3 protein; SEQ ID NO: 16.

Figure 20 is the amino acid sequence of BVH-3M protein; **SEQ ID** 20 NO: 55.

Figure 21 is the amino acid sequence of BVH-3AD protein; **SEQ ID** NO: 56.

25 Figure 22 is the amino acid sequence of L-BVH-3-AD protein; SEQ ID NO: 57.

Figure 23 is the amino acid sequence of NEW12 protein; SEQ ID NO: 58.

Figure 24 is the amino acid sequence of BVH-3C protein; SEQ ID NO: 59.

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Figure 25 is the amino acid sequence of BVH-11M protein; SEQ ID NO: 60.

5 Figure 26 is the amino acid sequence of BVH-11A protein; SEQ ID NO: 61.

Figure 27 is the amino acid sequence of BVH-11B (also called New13) protein; **SEQ ID NO: 62.**

Figure 28 is the amino acid sequence of BVH-11C protein; SEQ ID NO: 63.

Figure 29 is the amino acid sequence of NEW1 protein; SEQ ID NO: 64.

Figure 30 is the amino acid sequence of NEW2 protein; SEQ ID NO: 65.

20 Figure 31 is the amino acid sequence of NEW3 protein; **SEQ ID**NO: 66.

Figure 32 is the amino acid sequence of NEW4 protein; SEQ ID NO: 67.

Figure 33 is the amino acid sequence of NEW5 protein; SEQ ID NO: 68.

Figure 34 is the amino acid sequence of NEW6 protein; **SEQ ID** 30 NO: 69.

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Figure 35 is the amino acid sequence of NEW7 protein; SEQ ID NO: 70.

Figure 36 is the amino acid sequence of NEW8 protein; SEQ ID 5 NO: 71.

Figure 37 is the amino acid sequence of NEW9 protein; SEQ ID NO: 72.

10 Figure 38 is the amino acid sequence of BVH-11-2M protein; SEQ ID NO: 73.

Figure 39 is the amino acid sequence of NEW10 protein; SEQ ID NO: 74.

Figure 40 is the amino acid sequence of NEW11 protein; SEQ ID NO: 75.

Figure 41 is the DNA sequence of NEW12 gene; SEQ ID NO: 76.

Figure 42 is the amino acid sequence of NEW14 protein; SEQ ID NO: 77.

Figure 43 is the amino acid sequence of NEW15 protein; **SEQ ID** 25 NO: 78.

Figure 44 is the amino acid sequence of NEW16 protein; SEQ ID NO: 79.

30 Figure 45 is the DNA sequence of GBS BVH-71 gene; SEQ ID NO: 80.

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Figure 46 is the amino acid sequence of GBS BVH-71 protein; SEQ ID NO: 81.

Figure 47 is the DNA sequence of GAS BVH-71 gene; SEQ ID NO:82.

Figure 48 is the amino acid sequence of GAS BVH-71 protein; SEQ ID NO:83.

10 DETAILED DESCRIPTION OF THE INVENTION

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 95% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOs: 2, 4, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

30 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence

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chosen from SEQ ID NOs: 2, 4, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOs: 2, 4, 8, 10, 14, 16, 55 to 75, 77 to 79 or fragments, analogs or derivatives thereof.

10 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOs: 2, 8, 10, 16, 55, 56, 57, 58, 59, 64, 65, 66, 78 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOs: 2, 8, 10, 16, 55, 56, 57, 59, 64, 65, 66, 78 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOs: 4, 14, 58, 60, 61, 62, 63, 67, 68, 69, 70, 71, 72, 73, 74, 75, 77, 79 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOs: 4, 14, 60, 61, 62, 63, 67, 68, 69, 70,

71, 72, 73, 74, 75, 77, 79 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOs: 2, 4, 10, 14, 16, 55 to 75, 77 to 79 or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence chosen from SEQ ID NOs: 10, 55 to 75, 77, 78, 79 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence chosen from SEQ ID NOs: 55 to 75, 77, 78, 79 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOs: 2, 4, 6, 8, 10 or fragments, analogs or derivatives thereof.

30 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence

chosen from **SEQ ID NOs: 2, 4, 10, 14, 16** or fragments, analogs or derivatives thereof.

- According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOs: 2, 4, 14, 16 or fragments, analogs or derivatives thereof.
- 10 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID NO: 2 or fragments, analogs or derivatives thereof.
- 15 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID NO: 4 or fragments, analogs or derivatives thereof.
- According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID NO: 10 or fragments, analogs or derivatives thereof.
- 25 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID NO: 14 or fragments, analogs or derivatives thereof.
- 30 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID

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NO: 16 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID NO: 58 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID NO: 60 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID NO: 62 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID NO: 64 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID NO: 67 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID NO: 68 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID NO: 69 or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID NO: 72 or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID NO: 74 or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence SEQ ID NO: 77 or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from SEQ ID NOs: 2, 4, 6, 8, 10 or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen

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from SEQ ID NOs: 2, 4, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from SEQ ID NOs: 2, 4, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to
10 polypeptides characterized by the amino acid sequence chosen
from SEQ ID NOs: 2, 4, 8, 10, 14, 16, 55 to 75, 77 to 79 or
fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from SEQ ID NOs: 2, 4, 10, 14, 16, 55 to 75, 77 to 79 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to
20 polypeptides characterized by the amino acid sequence chosen
from SEQ ID NOs: 2, 4, 10, 14, 16 or fragments, analogs or
derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 2 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to 30 polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 4 or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 10 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 14 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 16 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from SEQ ID NOs: 10, 55 to 75, 77, 78, 79 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from SEQ ID NO: 10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from SEQ ID NO: 10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to

polypeptides characterized by the amino acid sequence chosen from SEQ ID NO: 10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives thereof.

- According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from SEQ ID NO: 10, 62, 64, 67, 68, 74, 77 or fragments, analogs or derivatives thereof.
- 10 According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 58 or fragments, analogs or derivatives thereof.
- 15 According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 62 or fragments, analogs or derivatives thereof.
- 20 According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 64 or fragments, analogs or derivatives thereof.
- 25 According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 67 or fragments, analogs or derivatives thereof.
- 30 According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 68 or fragments, analogs or

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derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 74 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 77 or fragments, analogs or derivatives thereof.

In a further embodiment, the present invention also relates to chimeric polypeptides which comprise one or more polypeptides or fragments, analogs or derivatives thereof as described in the present application.

In a further embodiment, the present invention also relates to chimeric polypeptides which comprise one or more polypeptides or fragments, analogs or derivatives thereof as defined in the figures of the present application.

In a further embodiment, the present application also relates to chimeric polypeptides which comprise two or more

25 polypeptides chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof; provided that the polypeptides or fragments, analogs or derivatives thereof are linked as to form a chimeric polypeptide.

In a further embodiment, the chimeric polypeptide will comprise two or more polypeptides chosen from SEQ ID NOs :10, 58, 60,

62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives thereof; provided that the polypeptides or fragments, analogs or derivatives thereof are linked as to form a chimeric polypeptide.

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In a further embodiment, the chimeric polypeptide will comprise two or more polypeptides chosen from SEQ ID NOs:10, 58, 60, 62, 64, 67, 68, 74, 77 or fragments, analogs or derivatives thereof; provided that the polypeptides or fragments, analogs or derivatives thereof are linked as to form a chimeric polypeptide.

In a further embodiment, the chimeric polypeptide will comprise two or more polypeptides chosen from SEQ ID NOs :10, 62, 64,

- 15 67, 68, 74, 77 or fragments, analogs or derivatives thereof; provided that the polypeptides or fragments, analogs or derivatives thereof are linked as to form a chimeric polypeptide.
- In a further embodiment, the chimeric polypeptide will comprise between 2 and 5 polypeptides.

In a further embodiment, the chimeric polypeptide will comprise between 2 and 4 polypeptides.

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In a further embodiment, the chimeric polypeptide will comprise between 2 and 3 polypeptides.

In a further embodiment, the chimeric polypeptide will comprise 30 2 polypeptides.

In a further embodiment, there is provided a chimeric polypeptide of formula (I):

 $A-(B)_{m}-(C)_{n}-D$ (I)

- 5 Wherein;
 - m is 0 or 1,
 - **n** is 0 or 1,
 - A is chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75,
 - 77 to 79, 81, 83 or fragments, analogs or derivatives thereof;
 B is chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75,
- 77 to 79, 81, 83 or fragments, analogs or derivatives thereof;
 - C is chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75,
 - 77 to 79, 81, 83 or fragments, analogs or derivatives thereof;
 - and

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15 D is chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75,
77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

In a further embodiment,

- A is chosen from SEQ ID NOs :10, 58, 60, 62, 64, 67, 68, 69,
- 72, 74, 77 or fragments, analogs or derivatives thereof;
- B is chosen from SEQ ID NOs :10, 58, 60, 62, 64, 67, 68, 69,
 - 72, 74, 77, or fragments, analogs or derivatives thereof;
 - C is chosen from SEQ ID NOs :10, 58, 60, 62, 64, 67, 68, 69,
 - 72, 74, 77 or fragments, analogs or derivatives thereof; and
- 25 D is chosen from SEQ ID NOs :10, 58, 60, 62, 64, 67, 68, 69,
 - 72, 74, 77 or fragments, analogs or derivatives thereof.

In a further embodiment,

- A is chosen from SEQ ID NOs :10, 58, 60, 62, 64, 67, 68, 74, 77
- 30 or fragments, analogs or derivatives thereof;
 - B is chosen from SEQ ID NOs :10, 58, 60, 62, 64, 67, 68, 74,
 - 77, or fragments, analogs or derivatives thereof;

C is chosen from SEQ ID NOs:10, 58, 60, 62, 64, 67, 68, 74, 77 or fragments, analogs or derivatives thereof; and D is chosen from SEQ ID NOs:10, 58, 60, 62, 64, 67, 68, 74, 77 or fragments, analogs or derivatives thereof.

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In one embodiment, chimeric polypeptides of the present invention comprise those wherein the following embodiments are present, either independently or in combination.

In a further embodiment, A is SEQ ID NOs :10, 58, 62, 64, 67,
68, 74, 77 or fragments, analogs or derivatives thereof.
In a further embodiment, A is SEQ ID NO :10 or fragments,
analogs or derivatives thereof.

In a further embodiment, A is SEQ ID NO :58 or fragments, analogs or derivatives thereof.

In a further embodiment, ${\bf A}$ is ${\bf SEQ}$ ${\bf ID}$ ${\bf NO}$:62 or fragments, analogs or derivatives thereof.

In a further embodiment, ${\bf A}$ is ${\bf SEQ}$ ${\bf ID}$ ${\bf NO}$:64 or fragments, analogs or derivatives thereof.

20 In a further embodiment, A is SEQ ID NO :67 or fragments, analogs or derivatives thereof.

In a further embodiment, A is SEQ ID NO :68 or fragments, analogs or derivatives thereof.

In a further embodiment, ${\bf A}$ is ${\bf SEQ}$ ${\bf ID}$ ${\bf NO}$:74 or fragments,

25 analogs or derivatives thereof.

In a further embodiment, ${\bf A}$ is **SEQ ID NO :77** or fragments, analogs or derivatives thereof.

In a further embodiment, B is SEQ ID NOs :10, 58, 62, 64, 67, 68, 74, 77 or fragments, analogs or derivatives thereof.

In a further embodiment, B is SEQ ID NO :10 or fragments, analogs or derivatives thereof.

In a further embodiment, B is SEQ ID NO :58 or fragments, analogs or derivatives thereof.

In a further embodiment, B is SEQ ID NO :64 or fragments, analogs or derivatives thereof.

5 In a further embodiment, B is SEQ ID NO :64 or fragments, analogs or derivatives thereof.

In a further embodiment, B is SEQ ID NO :67 or fragments, analogs or derivatives thereof.

In a further embodiment, ${\tt B}$ is ${\tt SEQ}$ ${\tt ID}$ ${\tt NO}$:68 or fragments,

10 analogs or derivatives thereof.

In a further embodiment, B is SEQ ID NO :74 or fragments, analogs or derivatives thereof.

In a further embodiment, B is SEQ ID NO: 77 or fragments, analogs or derivatives thereof.

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In a further embodiment, C is SEQ ID NOs :10, 58, 62, 64, 67, 68, 74, 77 or fragments, analogs or derivatives thereof.

In a further embodiment, C is SEQ ID NO :10 or fragments, analogs or derivatives thereof.

20 In a further embodiment, C is SEQ ID NO :58 or fragments, analogs or derivatives thereof.

In a further embodiment, ${\bf C}$ is ${\bf SEQ}$ ${\bf ID}$ ${\bf NO}$: 62 or fragments, analogs or derivatives thereof.

In a further embodiment, C is SEQ ID NO :64 or fragments,

25 analogs or derivatives thereof.

In a further embodiment, ${\bf C}$ is ${\bf SEQ}$ ${\bf ID}$ ${\bf NO}$: 67 or fragments, analogs or derivatives thereof.

In a further embodiment, C is SEQ ID NO: 68 or fragments, analogs or derivatives thereof.

30 In a further embodiment, C is SEQ ID NO: 74 or fragments, analogs or derivatives thereof.

In a further embodiment, C is SEQ ID NO: 77 or fragments,

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analogs or derivatives thereof.

In a further embodiment, D is SEQ ID NO :10, 58, 62, 64, 67, 68, 74, 77 or fragments, analogs or derivatives thereof.

5 In a further embodiment, **D** is **SEQ ID NO :10** or fragments, analogs or derivatives thereof.

In a further embodiment, ${\tt D}$ is ${\tt SEQ}$ ${\tt ID}$ ${\tt NO}$:58 or fragments, analogs or derivatives thereof.

In a further embodiment, ${\bf D}$ is ${\bf SEQ}$ ${\bf ID}$ ${\bf NO}$:62 or fragments,

10 analogs or derivatives thereof.

In a further embodiment, ${\tt D}$ is ${\tt SEQ}$ ${\tt ID}$ ${\tt NO}$:64 or fragments, analogs or derivatives thereof.

In a further embodiment, **D** is **SEQ ID NO :67** or fragments, analogs or derivatives thereof.

15 In a further embodiment, **D** is **SEQ ID NO :68** or fragments, analogs or derivatives thereof.

In a further embodiment, **D** is **SEQ ID NO :74** or fragments, analogs or derivatives thereof.

In a further embodiment, **D** is **SEQ ID NO :77** or fragments, analogs or derivatives thereof.

In a further embodiment, \mathbf{m} is 0. In a further embodiment, \mathbf{n} is 0.

In a further embodiment, \mathbf{m} and \mathbf{n} are 0.

In a further embodiment, m and n are 0, A is SEQ ID NO:64 or fragments, analogs or derivatives thereof, B is SEQ ID NO:62 or fragments, analogs or derivatives thereof.

In a further embodiment, m and n are 0, A is SEQ ID NO:62 or fragments, analogs or derivatives thereof, B is SEQ ID NO:64 or

fragments, analogs or derivatives thereof.

In accordance with the present invention, all nucleotides encoding polypeptides and chimeric polypeptides are within the scope of the present invention.

In a further embodiment, the polypeptides or chimeric polypeptides in accordance with the present invention are antigenic.

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In a further embodiment, the polypeptides or chimeric polypeptides in accordance with the present invention can elicit an immune response in an individual.

- In a further embodiment, the present invention also relates to polypeptides which are able to raise antibodies having binding specificity to the polypeptides or chimeric polypeptides of the present invention as defined above.
- An antibody that " has binding specificity" is an antibody that recognizes and binds the selected polypeptide but which does not substantially recognize and bind other molecules in a sample, e.g., a biological sample, which naturally includes the selected peptide. Specific binding can be measured using an ELISA assay in which the selected polypeptide is used as an antigen.

Unless otherwise defined, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs.

All publications, patent applications, patents, and other references mentioned herein are incorporated by reference in

their entirety. In case of conflict, the present specification, including definitions, will control. In addition, the materials, methods, and examples are illustrative only and not intended to be limiting.

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As used herein, "fragments", "derivatives" or "analogs" of the polypeptides of the invention include those polypeptides in which one or more of the amino acid residues are substituted with a conserved or non-conserved amino acid residue (preferably conserved) and which may be natural or unnatural. In one embodiment, derivatives and analogs of polypeptides of the invention will have about 70% identity with those sequences illustrated in the figures or fragments thereof. That is, 70% of the residues are the same. In a further embodiment, polypeptides will have greater than 75% homology. In a further embodiment, polypeptides will have greater than 80% homology. In a further embodiment, polypeptides will have greater than 85% homology. In a further embodiment, polypeptides will have greater than 90% homology. In a further embodiment, polypeptides will have greater than 95% homology. In a further embodiment, polypeptides will have greater than 99% homology. In a further embodiment, derivatives and analogs of polypeptides of the invention will have fewer than about 20 amino acid residue substitutions, modifications or deletions and more preferably less than 10. Preferred substitutions are those known in the art as conserved i.e. the substituted

30 In accordance with the present invention, polypeptides of the invention include both polypeptides and chimeric polypeptides.

residues share physical or chemical properties such as

hydrophobicity, size, charge or functional groups.

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Also included are polypeptides which have fused thereto other compounds which alter the polypeptides biological or pharmacological properties i.e. polyethylene glycol (PEG) to increase half-life; leader or secretory amino acid sequences for ease of purification; prepro- and pro- sequences; and (poly) saccharides.

Furthermore, in those situations where amino acid regions are found to be polymorphic, it may be desirable to vary one or more particular amino acids to more effectively mimic the different epitopes of the different streptococcus strains.

Moreover, the polypeptides of the present invention can be modified by terminal $-\mathrm{NH}_2$ acylation (eg. by acetylation, or thioglycolic acid amidation, terminal carbosy amidation, e.g. with ammonia or methylamine) to provide stability, increased hydrophobicity for linking or binding to a support or other molecule.

Also contemplated are hetero and homo polypeptide multimers of the polypeptide fragments, analogues and derivatives. These polymeric forms include, for example, one or more polypeptides that have been cross-linked with cross-linkers such as avidin/biotin, gluteraldehyde or dimethylsuperimidate. Such polymeric forms also include polypeptides containing two or more tandem or inverted contiguous sequences, produced from multicistronic mRNAs generated by recombinant DNA technology. Preferably, a fragment, analog or derivative of a polypeptide of the invention will comprise at least one antigenic region i.e. at least one epitope.

In order to achieve the formation of antigenic polymers (i.e.

synthetic multimers), polypeptides may be utilized having bishaloacetyl groups, nitroarylhalides, or the like, where the reagents being specific for thio groups. Therefore, the link between two mercapto groups of the different peptides may be a single bond or may be composed of a linking group of at least two, typically at least four, and not more than 16, but usually not more than about 14 carbon atoms.

In a particular embodiment, polypeptide fragments, analogs and derivatives of the invention do not contain a methionine (Met) starting residue. Preferably, polypeptides will not incorporate a leader or secretory sequence (signal sequence). The signal portion of a polypeptide of the invention may be determined according to established molecular biological techniques. In general, the polypeptide of interest may be isolated from a streptococcus culture and subsequently sequenced to determine the initial residue of the mature protein and therefore the sequence of the mature polypeptide.

20 According to another aspect, there are provided vaccine compositions comprising one or more streptococcus polypeptides of the invention in admixture with a pharmaceutically acceptable carrier diluent or adjuvant. Suitable adjuvants include oils i.e. Freund's complete or incomplete adjuvant; salts i.e. AlK(SO₄)₂, AlNa(SO₄)₂, AlNH₄(SO₄)₂, silica, kaolin, carbon polynucleotides i.e. poly IC and poly AU. Preferred adjuvants include QuilA and Alhydrogel. Vaccines of the invention may be administered parenterally by injection, rapid infusion, nasopharyngeal absorption, dermoabsorption, or bucal or oral. Pharmaceutically acceptable carriers also include tetanus toxoid.

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Vaccine compositions of the invention are used for the treatment or prophylaxis of streptococcus infection and/or diseases and symptoms mediated by streptococcus infection as described in P.R. Murray (Ed, in chief), E.J. Baron, M.A.

Pfaller, F.C. Tenover and R.H. Yolken. Manual of Clinical Microbiology, ASM Press, Washington, D.C. sixth edition, 1995, 1482p which are herein incorporated by reference. In one embodiment, vaccine compositions of the present invention are used for the treatment or prophylaxis of meningitis, otitis media, bacteremia or pneumonia. In one embodiment, vaccine compositions of the invention are used for the treatment or prophylaxis of streptococcus infection and/or diseases and symptoms mediated by streptococcus infection, in particular S.pneumoniae, group A streptococcus (pyogenes), group B streptococcus (GBS or agalactiae), dysgalactiae, uberis, nocardia as well as Staphylococcus aureus. In a further embodiment, the streptococcus infection is S.pneumoniae.

In a particular embodiment, vaccines are administered to those individuals at risk of streptococcus infection such as infants, elderly and immunocompromised individuals.

As used in the present application, the term " individuals" include mammals. In a further embodiment, the mammal is human.

Vaccine compositions are preferably in unit dosage form of about 0.001 to 100 $\mu g/kg$ (antigen/body weight) and more preferably 0.01 to 10 $\mu g/kg$ and most preferably 0.1 to 1 $\mu g/kg$ 1 to 3 times with an interval of about 1 to 6 week intervals between immunizations.

According to another aspect, there are provided polynucleotides encoding polypeptides characterized by the amino acid sequence chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

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In one embodiment, polynucleotides are those illustrated in SEQ ID Nos: 1, 3, 5, 7, 9, 11, 12, 13, 15, 76, 80, 82 which may include the open reading frames (ORF), encoding polypeptides of the invention. It will be appreciated that the polynucleotide sequences illustrated in the figures may be altered with degenerate codons yet still encode the polypeptides of the invention. Accordingly the present invention further provides polynucleotides which hybridize to the polynucleotide sequences herein above described (or the complement sequences thereof) having 50% identity between sequences. In one embodiment, at least 70% identity between sequences. In one embodiment, at least 75% identity between sequences. In one embodiment, at least 80% identity between sequences. In one embodiment, at least 85% identity between sequences. In one embodiment, at least 90% identity between sequences. In a further embodiment, polynucleotides are hybridizable under stringent conditions i.e. having at least 95% identity. In a further embodiment, more than 97% identity.

In a further embodiment, polynucleotides are those illustrated in SEQ ID NOs: 1, 3, 7, 9, 11, 12, 13, 15, 76, 80, 82 encoding polypeptides of the invention.

In a further embodiment, polynucleotides are those illustrated in SEQ ID NOs: 1, 3, 9, 11, 12, 13, 15, 76, 80, 82 which may include the open reading frames (ORF), encoding polypeptides of the invention.

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In a further embodiment, polynucleotides are those illustrated in SEQ ID NOs: 1, 3, 9, 11, 12, 13, 15, 76 which may include the open reading frames (ORF), encoding polypeptides of the invention.

In a further embodiment, polynucleotides are those illustrated in SEQ ID NOs: 1, 3, 7, 9, 11, 12, 13, 15, 76 which may include the open reading frames (ORF), encoding polypeptides of the invention.

In a further embodiment, polynucleotides are those illustrated in SEQ ID NOs: 1, 7, 9, 11, 15, 76 which may include the open reading frames (ORF), encoding polypeptides of the invention.

In a further embodiment, polynucleotides are those illustrated in SEQ ID NOs: 1, 9, 11, 15, 76 which may include the open reading frames (ORF), encoding polypeptides of the invention.

In a further embodiment, polynucleotides are those illustrated in SEQ ID NOs: 1, 7, 9, 11 which may include the open reading frames (ORF), encoding polypeptides of the invention.

In a further embodiment, polynucleotides are those illustrated in SEQ ID NO: 1, encoding polypeptides of the invention.

In a further embodiment, polynucleotides are those illustrated in SEQ ID NO :7, encoding polypeptides of the invention.

In a further embodiment, polynucleotides are those illustrated in SEQ ID NO :9, encoding polypeptides of the invention.

In a further embodiment, polynucleotides are those illustrated in SEQ ID NO :11, encoding polypeptides of the invention.

In a further embodiment, polynucleotides are those illustrated in SEQ ID NO :15, encoding polypeptides of the invention.

In a further embodiment, polynucleotides are those illustrated in **SEQ ID NOs**: 3, 12, 13, 76, encoding polypeptides of the invention.

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In a further embodiment, polynucleotides are those illustrated in SEQ ID NO :3, encoding polypeptides of the invention.

In a further embodiment, polynucleotides are those illustrated in SEQ ID NO :12, encoding polypeptides of the invention.

In a further embodiment, polynucleotides are those illustrated in SEQ ID NO :13, encoding polypeptides of the invention.

20 In a further embodiment, polynucleotides are those illustrated in **SEQ ID NO :76**, encoding polypeptides of the invention.

As will be readily appreciated by one skilled in the art, polynucleotides include both DNA and RNA.

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The present invention also includes polynucleotides complementary to the polynucleotides described in the present application.

In a further aspect, polynucleotides encoding polypeptides of the invention, or fragments, analogs or derivatives thereof, may be used in a DNA immunization method. That is, they can be

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incorporated into a vector which is replicable and expressible upon injection thereby producing the antigenic polypeptide in vivo. For example polynucleotides may be incorporated into a plasmid vector under the control of the CMV promoter which is functional in eukaryotic cells. Preferably the vector is injected intramuscularly.

According to another aspect, there is provided a process for producing polypeptides of the invention by recombinant techniques by expressing a polynucleotide encoding said polypeptide in a host cell and recovering the expressed polypeptide product. Alternatively, the polypeptides can be produced according to established synthetic chemical techniques i.e. solution phase or solid phase synthesis of oligopeptides which are ligated to produce the full polypeptide (block ligation).

General methods for obtention and evaluation of
polynucleotides and polypeptides are described in the following
references: Sambrook et al, Molecular Cloning: A Laboratory
Manual, 2nd ed, Cold Spring Harbor, N.Y., 1989; Current
Protocols in Molecular Biology, Edited by Ausubel F.M. et al.,
John Wiley and Sons, Inc. New York; PCR Cloning Protocols, from
Molecular Cloning to Genetic Engineering, Edited by White B.A.,
Humana Press, Totowa, New Jersey, 1997, 490 pages; Protein
Purification, Principles and Practices, Scopes R.K., SpringerVerlag, New York, 3rd Edition, 1993, 380 pages; Current
Protocols in Immunology, Edited by Coligan J.E. et al., John
Wiley & Sons Inc., New York which are herein incorporated by
reference.

For recombinant production, host cells are transfected with

vectors which encode the polypeptide, and then cultured in a nutrient media modified as appropriate for activating promoters, selecting transformants or amplifying the genes. Suitable vectors are those that are viable and replicable in the chosen host and include chromosomal, non-chromosomal and 5 synthetic DNA sequences e.g. bacterial plasmids, phage DNA, baculovirus, yeast plasmids, vectors derived from combinations of plasmids and phage DNA. The polypeptide sequence may be incorporated in the vector at the appropriate site using restriction enzymes such that it is operably linked to an 10 expression control region comprising a promoter, ribosome binding site (consensus region or Shine-Dalgarno sequence), and optionally an operator (control element). One can select individual components of the expression control region that are appropriate for a given host and vector according to 15 established molecular biology principles (Sambrook et al, Molecular Cloning: A Laboratory Manual, 2nd ed, Cold Spring Harbor, N.Y., 1989; Current Protocols in Molecular Biology, Edited by Ausubel F.M. et al., John Wiley and Sons, Inc. New York incorporated herein by reference). Suitable promoters 20 include but are not limited to LTR or SV40 promoter, E.coli lac, tac or trp promoters and the phage lambda P_{L} promoter. Vectors will preferably incorporate an origin of replication as well as selection markers i.e. ampicilin resistance gene. Suitable bacterial vectors include pET, pQE70, pQE60, pQE-9, 25 pbs, pD10 phagescript, psiX174, pbluescript SK, pbsks, pNH8A, pNH16a, pNH18A, pNH46A, ptrc99a, pKK223-3, pKK233-3, pDR540, pRIT5 and eukaryotic vectors pBlueBacIII, pWLNEO, pSV2CAT, pOG44, pXT1, pSG, pSVK3, pBPV, pMSG and pSVL. Host cells may be bacterial i.e. <u>E.coli</u>, <u>Bacillus subtilis</u>, <u>Streptomyces</u>; 30 fungal i.e. Aspergillus niger, Aspergillus nidulins; yeast i.e.

<u>Saccharomyces</u> or eukaryotic i.e. CHO, COS.

Upon expression of the polypeptide in culture, cells are typically harvested by centrifugation then disrupted by physical or chemical means (if the expressed polypeptide is not secreted into the media) and the resulting crude extract retained to isolate the polypeptide of interest. Purification of the polypeptide from culture media or lysate may be achieved by established techniques depending on the properties of the polypeptide i.e. using ammonium sulfate or ethanol

10 precipitation, acid extraction, anion or cation exchange chromatography, phosphocellulose chromatography, hydrophobic interaction chromatography, hydroxylapatite chromatography and lectin chromatography. Final purification may be achieved using HPLC.

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The polypeptide may be expressed with or without a leader or secretion sequence. In the former case the leader may be removed using post-translational processing (see US 4,431,739; US 4,425,437; and US 4,338,397 incorporated herein by reference) or be chemically removed subsequent to purifying the expressed polypeptide.

According to a further aspect, the streptococcus polypeptides of the invention may be used in a diagnostic test for streptococcus infection, in particular <u>S. pneumoniae</u> infection. Several diagnostic methods are possible, for example detecting streptococcus organism in a biological sample, the following procedure may be followed:

- a) obtaining a biological sample from a patient;
- 30 b) incubating an antibody or fragment thereof reactive with a streptococcus polypeptide of the invention with the biological sample to form a mixture; and

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c) detecting specifically bound antibody or bound fragment in the mixture which indicates the presence of streptococcus.

Alternatively, a method for the detection of antibody specific to a streptococcus antigen in a biological sample containing or suspected of containing said antibody may be performed as follows:

- a) obtaining a biological sample from a patient;
- b) incubating one or more streptococcus polypeptides of the invention or fragments thereof with the biological sample to form a mixture; and
 - c) detecting specifically bound antigen or bound fragment in the mixture which indicates the presence of antibody specific to streptococcus.

One of skill in the art will recognize that this diagnostic test may take several forms, including an immunological test such as an enzyme-linked immunosorbent assay (ELISA), a radioimmunoassay or a latex agglutination assay, essentially to determine whether antibodies specific for the protein are present in an organism.

The DNA sequences encoding polypeptides of the invention may also be used to design DNA probes for use in detecting the presence of streptococcus in a biological sample suspected of containing such bacteria. The detection method of this invention comprises:

- a) obtaining the biological sample from a patient;
- b) incubating one or more DNA probes having a DNA sequence
 encoding a polypeptide of the invention or fragments
 thereof with the biological sample to form a mixture; and
 - c) detecting specifically bound DNA probe in the mixture

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which indicates the presence of streptococcus bacteria.

The DNA probes of this invention may also be used for detecting circulating streptococcus i.e. S.pneumoniaenucleic acids in a sample, for example using a polymerase chain reaction, as a method of diagnosing streptococcus infections. The probe may be synthesized using conventional techniques and may be immobilized on a solid phase, or may be labelled with a detectable label. A preferred DNA probe for this application is an oligomer having a sequence complementary to at least about 6 contiguous nucleotides of the streptococcus pneumoniae polypeptides of the invention.

Another diagnostic method for the detection of streptococcus in a patient comprises:

- a) labelling an antibody reactive with a polypeptide of the invention or fragment thereof with a detectable label;
- b) administering the labelled antibody or labelled fragment to the patient; and
- 20 c) detecting specifically bound labelled antibody or labelled fragment in the patient which indicates the presence of streptococcus.

A further aspect of the invention is the use of the streptococcus polypeptides of the invention as immunogens for the production of specific antibodies for the diagnosis and in particular the treatment of streptococcus infection. Suitable antibodies may be determined using appropriate screening methods, for example by measuring the ability of a particular antibody to passively protect against streptococcus infection in a test model. One example of an animal model is the mouse model described in the examples herein. The antibody may be a

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whole antibody or an antigen-binding fragment thereof and may belong to any immunoglobulin class. The antibody or fragment may be of animal origin, specifically of mammalian origin and more specifically of murine, rat or human origin. It may be a natural antibody or a fragment thereof, or if desired, a recombinant antibody or antibody fragment. The term recombinant antibody or antibody fragment means antibody or antibody fragment which was produced using molecular biology techniques. The antibody or antibody fragments may be polyclonal, or preferably monoclonal. It may be specific for a number of epitopes associated with the streptococcus pneumoniae

polypeptides but is preferably specific for one.

Without limiting its scope, the present invention also relates to new antigens designated BVH-3, BVH-11, BVH-11-2, BVH-28 and BVH-71. The present invention also relates to truncated polypeptides comprising fragments of the new antigens designated BVH-3, BVH-11, BVH-11-2, BVH-28 and BVH-71. The present invention also relates to chimeric polypeptides comprising fragments of the new antigens designated BVH-3, BVH-11, BVH-11-2, BVH-28 and BVH-71. The following is a reference table summarizing the relation between the antigens of the present invention:

Family	Nucleotide SEQ ID	Polypeptide SEQ ID NO
BVH-3		
BVH-3	1, 11	2
BVH-3A	7	8
BVH-3B	9	10
BVH-3 SP63	15	16
BVH-3M		55
BVH-3AD		56
L-BVH-3AD		57
New12	76	58
BVH-3C		59
New1		64
New2		65
New3		66
New15		78
BVH-11		
BVH-11	3, 12	4
BVH-11-2	13	14
BVH-11M		60
BVH-11A		61
BVH-11B also		62
referred to as		
NEW13		
BVH-11C		63
New4		67
New5		68
New6		69
New7		70
New8		71
New9		72
BVH-11-2M		73
New10		74
New11		75
New12	76	58
New14		77
New16		79
BVH-28		
BVH-28	5	6
BVH-71		
GBS	80	81
GAS	82	83

EXAMPLE 1

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This example illustrates the cloning of S. pneumoniae genes.

The coding region of \underline{S} . $\underline{pneumoniae}$ gene BVH-3 (SEQ ID NO: 1) and the coding region of \underline{S} . $\underline{pneumoniae}$ gene BVH-28 (SEQ ID NO: 5) were amplified by PCR (DNA Thermal Cycler GeneAmp PCR system 2400 Perkin Elmer, San Jose, CA) from genomic DNA of serogroup 6 S. pneumoniae strain SP64 using the oligos that contained base 10 extensions for the addition of restriction sites BglII (AGATCT) and XbaI (TCTAGA). PCR products were purified from agarose gel using a QIAquick gel extraction kit from QIAgen (Chatsworth, CA), digested BglII-XbaI (Pharmacia Canada Inc, Baie d'Urfé, Canada), extracted with phenol : chloroform and precipitated 15 with ethanol. The Superlinker vector pSL301 (Invitrogen, San Diego, CA) was digested with BglII and XbaI and purified from agarose gel using a QIAquick gel extraction kit from QIAgen (Chatsworth, CA). The BglII-XbaI genomic DNA fragments were ligated to the BglII-XbaI pSL301 vector. The ligated products 20 were transformed into <u>E. coli</u> strain DH5a [f80 lacZ DM15 endA1 recA1 hsdR17 ($^{r}K^{-m}K^{+}$) supE44 thi-11 gyrA96 relA1 D(lacZYAargF)U169] (Gibco BRL, Gaithersburg, MD) according to the method of Simanis (Hanahan, D. DNA Cloning, 1985, D.M. Glover (ed), pp. 109-135). Recombinant pSL301 plasmids (rpSL301) containing 25 either BVH-3 or BVH-28 gene were purified using a QIAgen kit (Chatsworth, CA) and DNA inserts were confirmed by nucleotide sequence analysis (Taq Dye Deoxy Terminator Cycle Sequencing kit, ABI, Foster City, CA). Recombinant rpSL301 (rpSL301) were digested with the restriction enzymes BglII (AGATCT) and XhoI 30

QIAquick gel extraction kit from QIAgen (Chatsworth, CA). pET-

(CTCGAG). DNA fragments BglII-XhoI were purified using the

32c(+) expression vector (Novagen, Madison, WI) containing the thioredoxin-His·Tag sequence was digested with BamHI (GGATCC) and XhoI and gel extracted using the QIAquick gel extraction kit from QIAgen (Chatsworth, CA). The BglII-XhoI DNA fragments were ligated to the BamHI-XhoI pET-32c(+) vector to create the coding 5 sequence for thioredoxin-His·Tag-BVH-3 or thioredoxin-His·Tag-BVH-28 fusion protein. The ligated products were transformed into <u>E. coli</u> strain DH5a [f80 lacZ DM15 endA1 recA1 hsdR17 (r K $^{-}$ $^{m}K^{+}$) supE44 $thi-1l^{-}$ gyrA96 relA1 D(lacZYA-argF)U169] (Gibco BRL, Gaithersburg, MD) according to the method of Simanis (Hanahan, 10 D. DNA Cloning, 1985, D.M. Glover (ed), pp. 109-135). Recombinant pET-32c(+) plasmids were purified using a QIAgen kit (Chatsworth, CA) and the nucleotide sequences at the fusion sites of thioredoxin-His·Tag and DNA insert were verified by DNA sequencing (Tag Dye Deoxy Terminator Cycle Sequencing kit, ABI, 15

EXAMPLE 2

Foster City, CA).

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This example illustrates the cloning of <u>S. pneumoniae</u> protein genes in CMV plasmid pCMV-GH.

The DNA coding region of a <u>S. pneumoniae</u> protein was inserted in phase downstream of a human growth hormone (hGH) gene which was under the transcriptional control of the cytomegalavirus (CMV) promotor in the plasmid vector pCMV-GH (Tang et al., Nature, 1992, 356:152). The CMV promotor is non functional plasmid in <u>E. coli</u> cells but active upon administration of the plasmid in eukaryotic cells. The vector also incorporated the ampicillin resistance gene.

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The coding region of BVH-3 gene (SEQ ID NO: 1) and BVH-28 gene (SEQ ID NO: 5) were obtained from rpSL301 (see example 1) using restriction enzymes BglII (AGATCT) and XbaI (TCTAGA). digested products were purified from agarose gel using the QIAquick gel extraction kit from QIAgen (Chatsworth, CA). 5 pCMV-GH vector (Laboratory of Dr. Stephen A. Johnston, Department of Biochemistry, The University of Texas, Dallas, Texas) containing the human growth hormone to create fusion proteins was digested with BglII and XbaI and purified from agarose gel using the QIAquick gel extraction kit from QIAgen 10 (Chatsworth, CA). The BglII-XbaI DNA fragments were ligated to the BglII-XbaI pCMV-GH vector to create the hGH-BVH-3 or hGH-BVH-28 fusion protein under the control of the CMV promoter. The ligated products were transformed into E. coli strain DH5a[f80 lacZ DM15 endA1 recA1 hsdR17 ($^{r}K^{-m}K^{+}$) supE44 thi-11 gyrA96 relA1 15 D(lacZYA-argF)U169] (Gibco BRL, Gaithersburg, MD) according to the method of Simanis (Hanahan, D. DNA Cloning, 1985, D.M. Glover (ed), pp. 109-135). The recombinant pCMV plasmids were purified using a QIAgen kit (QIAgen, Chatsworth, CA).

The coding region of BVH-11 gene (SEQ ID NO: 3) was amplified by PCR (DNA Thermal Cycler GeneAmp PCR system 2400 Perkin Elmer, San Jose, CA) from genomic DNA of serogroup 6 S. pneumoniae strain SP64 using the oligos that contained base extensions for the addition of restriction sites BglII (AGATCT) and HindIII (AAGCTT). The PCR product was purified from agarose gel using a QIAquick gel extraction kit from QIAgen (Chatsworth, CA), digested with restriction enzymes (Pharmacia Canada Inc, Baie d'Urfe, Canada), extracted with phenol: chloroform and precipitated with ethanol. The pCMV-GH vector (Laboratory of Dr. Stephen A. Johnston, Department of Biochemistry, The University of Texas, Dallas, Texas) was digested with BglII and

HindIII and purified from agarose gel using the QIAquick gel extraction kit from QIAgen (Chatsworth, CA). The BglII-HindIII DNA fragment was ligated to the BglII-HindIII pCMV-GH vector to create the hGH-BVH-11 fusion protein under the control of the 5 CMV promoter. The ligated products were transformed into <u>E. colistarian DH5a[f80 lacZ DM15 endA1 recA1 hsdR17 (*K-mK+) supE44 thi-11-gyrA96 relA1 D(lacZYA-argF)U169] (Gibco BRL, Gaithersburg, MD) according to the method of Simanis (Hanahan, D. DNA Cloning, 1985, D.M. Glover (ed), pp. 109-135). The recombinant pCMV plasmid was purified using a QIAgen kit (Chatsworth, CA) and the nucleotide sequence of the DNA insert was verified by DNA sequencing.</u>

15 EXAMPLE 3

This example illustrates the use of DNA to elicit an immune response to S. pneumoniae antigens.

20 A group of 8 female BALB/c mice (Charles River, St-Constant, Québec, Canada) were immunized by intramuscular injection of 50 μl three times at two- or three-week intervals with 100 μg of recombinant pCMV-GH encoding the BVH-3, BVH-11 or the BVH-28 gene in presence of 50 μg of granulocyte-macrophage colony25 stimulating factor (GM-CSF) - expressing plasmid pCMV-GH-GM-CSF (Laboratory of Dr. Stephen A. Johnston, Department of Biochemistry, The University of Texas, Dallas, Texas). As control, a group of mice were injected with 100 μg of pCMV-GH in presence of 50 μg of pCMV-GH-GM-CSF. Blood samples were collected from the orbital prior to each immunization and seven days following the third injection and serum antibody responses

were determined by ELISA using thioredoxin-His·Tag-S. pneumoniae fusion protein as coating antigen. DNA immunization with recombinant plasmid pCMV-GH encoding the BVH-3, BVH-11 or the BVH-28 S. pneumoniae protein induced antibody reactive against the respective recombinant protein. The reciprocal antibody titers, defined as the highest serum dilution at which the absorbance values were 0.1 above the background values, were above 4×10^3 .

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EXAMPLE 4

This example illustrates the production and purification of recombinant <u>S. pneumoniae</u> proteins.

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The recombinant pET plasmids containing the BVH-3, BVH-11 or the BVH-28 gene corresponding to the SEQ ID NO: 1 , SEQ ID NO: 3 or the SEQ ID NO: 5 respectively were transformed by electroporation (Gene Pulser II apparatus, BIO-RAD Labs, 20 Mississauga, Canada) into E. coli strain AD494 (DE3) (Dara leu7697 DlacX74 DphoA PvuII phoR DmalF3 F'[lac+(lacIq) pro] trxB::Kan) (Novagen, Madison, WI). In this strain of E. coli, the T7 promotor controlling expression of the fusion protein is specifically recognized by the T7 RNA polymerase (present on the 25 1DE3 prophage) whose gene is under the control of the lac promotor which is inducible by isopropyl-ß-d-thiogalactopyranoside (IPTG). The transformant AD494(DE3)/rpET was grown at 37°C with agitation at 250 rpm in LB broth (peptone 10g/L, yeast extract 5g/L, NaCl 10g/L) containing 100μg of 30 ampicillin (Sigma-Aldrich Canada Ltd., Oakville, Canada) per ml until the A_{600} reached a value of 0.6. In order to induce the production of the thioredoxin-His·Tag-BVH-3, thioredoxin-

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His·Tag-BVH-11 or thioredoxin-His·Tag-BVH-28 fusion protein, the cells were incubated for 2 additional hours in the presence of IPTG at a final concentration of 1 mM. Induced cells from a 100 ml culture were pelleted by centrifugation and frozen at -70°C.

The purification of the fusion proteins from the soluble cytoplasmic fraction of IPTG-induced AD494 (DE3) /rpET was done by affinity chromatography based on the properties of the His-Tag sequence (6 consecutive histidine residues) to bind to divalent cations (Ni^{2+}) immobilized on the His·Bind metal chelation resin. Briefly, the pelleted cells obtained from a 100mL culture induced with IPTG were resuspended in phosphate-buffered (PBS):500mM NaCl pH7.1, sonicated and spun at 20,000 X g for 20 min to remove debris. The supernatant was filtered (0.22μm pore size membrane) and deposited on a HiTrap® 1mL chelating prepacked ready-to-use column (Pharmacia Biotech, Baie d'Urfé, Canada). The thioredoxin-His·Taq-S. pneumoniae fusion protein was eluted with 1M imidazole-500mM NaCl-PBS pH7.1. The removal of the salt and imidazole from the sample was done by dialysis against PBS at 4°C. The quantities of fusion protein obtained from the soluble fraction of E. coli was estimated by MicroBCA (Pierce, Rockford, Illinois).

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EXAMPLE 5

This example illustrates the protection of mice against fatal pneumococcal infection by immunization.

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Groups of 8 female BALB/c mice (Charles River) were immunized subcutaneously three times at three-week intervals with either

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25 μg of affinity purified thioredoxin-His·Tag-BVH-3 fusion protein in presence of 15 μg of QuilA adjuvant (Cedarlane Laboratories Ltd, Hornby, Canada) or, as control, with QuilA adjuvant alone in PBS. Blood samples were collected from the orbital sinus on day 1, 22 and 43 prior to each immunization and seven days (day 50) following the third injection. One week later the mice were challenged with approximately 10⁶ CFU of the type 3 <u>S. pneumoniae</u> strain WU2. Samples of the <u>S. pneumoniae</u> challenge inoculum were plated on chocolate agar plates to determine the CFU and to verify the challenge dose. Deaths were recorded for a period of 14 days and on day 14 post-challenge, the surviving mice were sacrificied and blood samples tested for the presence of <u>S. pneumoniae</u> organisms. The survival data are shown in table 1.

Prechallenge sera were analyzed for the presence of antibodies reactive with <u>S. pneumoniae</u> by standard immunoassays. Elisa and immunoblot analyses indicated that immunization with recombinant <u>S. pneumoniae</u> protein produced in <u>E. coli</u> elicited antibodies reactive with both, recombinant and native pneumococcal protein.

Table 1. Protection mediated by recombinant BVH-3 protein

Immunogen	No. of mice alive : no. of mice	Median day of
	dead	death
	14 days post-challenge	
BVH-3	8:0	>14
none	0:8	1

25 All mice immunized with BVH-3 recombinant protein survived to infection while none of the control mice given adjuvant alone

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survived. There was a significant difference in survival between the two groups of mice (P<0.0001, log rank test for nonparametric analysis of survival curves; P=0.0002, Fisher's exact test). All hemocultures from surviving mice were negative at day 14 post-challenge.

EXAMPLE 6

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This example describes the cloning of $\underline{BVH-3}$ and $\underline{BVH-11}$ genes from a variety of \underline{S} . pneumoniae strains and the molecular conservation of these genes.

Molecular analysis of chromosomal DNA from various <u>S.</u>
<u>pneumoniae</u> isolates with DNA probes spanning different regions of <u>BVH-3</u> or <u>BVH-11</u> revealed the presence of one <u>BVH-3</u> gene copy and two <u>BVH-11</u> gene copies. The two <u>BVH-11</u> gene copies are not identical and the genes were arbitrarily designated <u>BVH-11</u> (SEQ ID NO:12; ORF at nucleotides 45 to 2567) and <u>BVH-11-2</u> (SEQ ID NO:13; ORF at nucleotides 114 to 2630).

The first amino acids of the BVH-3 and BVH-11 coding regions have the characteristics of leader sequences also known as signal peptides. The consensus signal peptidase cleavage site L-X-X-C of lipoprotein modification/processing sites was present in the sequences. Mature BVH-3, BVH-11 and BVH-11-2 proteins from <u>S. pneumoniae</u> SP64 have 1019, 821 and 819 amino acids, respectively. The regions of <u>S. pneumoniae</u> genes coding for mature BVH-3, termed BVH-3M, (nucleotides 1837 - 4896; SEQ. ID. NO: 11), BVH-11M (nucleotides 102-2567; SEQ. ID. NO: 12) and BVH-11-2M (nucleotides 171-2630; SEQ. ID. NO: 13),

were amplified by PCR (DNA Thermal Cycler GeneAmp PCR system 2400 Perkin Elmer, San Jose, CA) from genomic DNA of 6 or 7 S. pneumoniae strains. Serogroup 6 S. pneumoniae SP64 and serogroup 9 SP63 clinical isolates were provided by the laboratoire de la santé publique du Québec, Sainte-Anne-de-Bellevue; serotype 4 strain JNR.7/87 was provided by Andrew Camilli, Tufts University School of Medicine, Boston; Rx1 strain, a nonencapsulated derivative of the type 2 strain D39 and the type 3 strains A66 and WU2 were provided by David E. 10 Briles from University of Alabama, Birmingham and the type 3 clinical isolate P4241 was provided by the centre de recherche en infectiologie du centre hospitalier de l'université Laval, Sainte-Foy. The sets of oligonucleotide primers OCRR479-OCRR480; HAMJ160-OCRR488 and HAMJ160-HAMJ186, that contained base extensions for the addition of restriction sites were used 15 for the amplification of BVH-3, BVH-11 and BVH-11-2 gene, respectively, with the exception of BVH-11 gene from SP64 strain which was amplified using the set of primers consisting of HAMJ487 and OCRR488. Primer sequences are listed below PCR products were purified from agarose gel using 20 a QIAquick gel extraction kit from QIAgen (Chatsworth, CA) and digested BglII-XbaI or BglII-HindIII (Pharmacia Canada Inc, Baie d'Urfé, Canada). Digestions were cleaned using a QIAquick PCR purification kit from QIAgen (Chatsworth, CA). The PCR 25 products were ligated to the BglII-XbaI or BglII-HindIII pSL301 vector. The ligated products were transformed into E. <u>coli</u> strain DH5 α [ϕ 80 *lac*Z Δ M15 *end*A1 *rec*A1 *hsd*R17 ($^{r}K^{-m}K^{+}$) $supE44 thi-1\lambda^{-} qyrA96 relA1 \Delta(lacZYA-argF)U169]$ (Gibco BRL, Gaithersburg, MD) according to the method of Simanis (Hanahan,

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Recombinant pSL301 plasmids (rpSL301) containing <u>BVH-3</u>, <u>BVH-11</u>

D. DNA Cloning, 1985, D.M. Glover (ed), pp. 109-135).

or <u>BVH11-2</u> were purified using a QIAgen kit (Chatsworth, CA) and DNA inserts were sequenced (Taq Dye Deoxy Terminator Cycle Sequencing kit, ABI, Foster City, CA). The figures 11 and 12 depict the consensus sequence established from the BVH-3, and BVH-11 deduced amino acid sequences, respectively. Comparison 5 of BVH-3 protein sequences revealed 99 to 100% identity of sequences for all strains with the exception that BVH-3 from serogroup 9 SP63 strain (SEQ. ID. NO: 15 and SEQ. ID. NO: 16) misses a stretch of 177 amino acids corresponding to residues 244 to 420 on BVH-3 protein sequence of S. pneumoniae SP64. 10 Analysis of sequences of additional serogroup 9 strains revealed BVH-3 molecule having the same deletion in 3 out of 4 strains thus suggesting that the 3 strains are members of a \underline{S} . pneumoniae serogroup 9 clone.

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Comparison of 13 BVH-11 nucleotide sequences obtained from 7 S. pneumoniae strains, revealed that the nucleotide sequences are very similar. Computer analysis (MacVector, Clustal W 1.4) using multiple alignment of the predicted BVH-11 protein sequences revealed that these sequences were 75% identical and 20 82 % homologous on a length of 834 amino acids. Pairwise alignment revealed 80 to 100% identity (Figure 13). sequences showed great similarity in overall organization. Variability in the primary sequence of these proteins is almost restricted to the last 125 amino acids in the C-terminal 25 portion of the proteins. This region constitutes a domain. Close examination of this domain revealed two groups of sequences. The first 9 sequences from the figure 13 belong to one group while the last 4 sequences belong to another group. A 39% identity value is obtained when the domain sequences of the 30 13 proteins are compared (MacVector, Clustal W 1.4).

identity value increased to more than 92% when sequences belonging to a same group are compared.

5 EXAMPLE 7

This example illustrates the homology of portions of $\underline{BVH-3}$ and $\underline{BVH-11}$ genes.

10 Molecular analysis with DNA probes derived from BVH-3 and BVH-11 genes indicated that BVH-3 and BVH-11 were related. blot hybridization studies, DNA probe consisting of either, BVH-3 or BVH-11, gene sequence hybridized to both, BVH-3 and BVH-11 genes thus indicating that BVH-3 and BVH-11 genes shared homologous sequences. Comparison of sequences revealed that 15 the ORFs and the proteins were 43 and 33% identical, respectively. Closer examination revealed that the region corresponding to amino acids 1 to 225 in BVH-3 and 1 to 228 in BVH-11 were 73 and 75% identical at the DNA and protein level, 20 respectively. In contrast, the 3' regions corresponding to amino acids 226 to 1039 from BVH-3 and amino acids 229-840 from BVH-11 were only 34 and 22% identical at the DNA and protein level, respectively. Thus the 5' termini of BVH-3 and BVH-11 genes appear to contain highly conserved sequences while the 25 remaining parts of the genes are highly divergent. results suggest that BVH-3 and BVH-11 might share similar functions mediated by sequences present in the conserved region whereas BVH-3- and BVH-11-specific functions might be mediated by sequences in the divergent region.

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This example describes the cloning of truncated $\underline{BVH-3}$, $\underline{BVH-11}$ and $\underline{BVH-11-2}$ genes by polymerase chain reaction (PCR) and the expression of truncated $\underline{BVH-3}$ and $\underline{BVH-11}$ molecules.

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Gene fragments were amplified by PCR using pairs of oligonucleotide engineered to amplify fragments spanning the BVH-3 (SEQ ID NO: 1 and SEQ ID NO: 11), BVH-11 (SEQ ID NO: 3 and SEQ ID NO: 12) or BVH-11-2 (SEQ ID NO: 13) gene from S. pneumoniae strain SP64. Each of the primers had a restriction endonuclease site at the 5' end, thereby allowing directional in-frame cloning of the amplified product into the digested plasmid vector (Tables 2 and 3). PCR-amplified products were digested with restriction endonucleases and ligated to either linearized plasmid pSL301 (see example 1), pCMV-GH (see example 2) or pET (Novagen, Madison, WI) expression vector digested likewise or digested with enzymes that produce compatible cohesive ends. Recombinant pSL301 and recombinant pCMV-GH plasmids were digested with restriction enzymes for the inframe cloning in pET expression vector. Clones were first stabilized in E. coli DH5 α before introduction into E. coliBL21(λ DE3) or AD494 (λ DE3) for expression of truncated BVH-3 or BVH-11 molecules. Each of the resultant plasmid constructs was confirmed by nucleotide sequence analysis. The recombinant proteins were expressed as N-terminal fusions with the thioredoxin and His-tag or as C-terminal fusions with an His-The expressed recombinant proteins were purified from supernatant fractions obtained from centrifugation of sonicated IPTG-induced <u>E. coli</u> cultures using a His-Bind metal chelation resin (QIAgen, Chatsworth, CA). The gene products generated are listed in the table 3. The gene products corresponding to

the N-terminal region including the signal sequence are designated as Lipidated-proteins or lipoproteins (L-proteins). The gene products corresponding to the N-terminal region lacking the signal sequence are identified as protein without signal sequence (w/o ss).

Table 2. List of PCR oligonucleotide primers

l	l	

Primer	SEQ.	Sequence 5' - 3'	Nucleotide	Restric-
	ID.		position	tion
				sites
OCRR 479	17	cagtagatetgtgeetatgeactaaac	SEQ ID 1:61-	BglII
			78	
OCRR 480	18	gatetetagaetaetgetatteettaegetatg	SEQ	XbaI
			ID 11 :4909-	
			4887	
OCRR 497	19	atcactcgagcattacctggataatcctgt	SEQ	XhoI
			ID 1 :1525-	
			1506	
OCRR 498	20	ctgctaagcttatgaaagatttagat	SEQ	HindIII
			ID 1 :1534-	
			1548	
OCRR 499	21	gatactcgagctgctattccttac	SEQ	XhoI
			ID 11 :4906-	
			4893	
HAMJ 172	22	gaatetegagttaagetgetgetaatte	SEQ ID 1:	XhoI
			675-661	
HAMJ 247	23	gacgetegagegetatgaaateagataaatte	SEQ ID	XhoI
			1:3117-3096	
HAMJ 248	24	gacgetegagggeattacetggataateetgtteatg	SEQ ID	XhoI
			1:1527-1501	
HAMJ 249	25	cagtagatetetteateatttattgaaaagagg	SEQ ID 11:	BglII
121 2020 200			1749-1771	
HAMJ 278	26	ttatttetteeatatggaettgaeagaagageaaattaag	SEQ ID	NdeI
			1:1414-1437	
HAMJ 279	27	cgccaagettegetatgaaatcagataaatte	SEQ ID	HindIII
11/1/11/2/7/	1	190000000000000000000000000000000000000	1:3117-3096	
HAMJ 280	28	cgccaagcttttccacaatataagtcgattgatt	SEQ ID	HindIII
111111111111111111111111111111111111111		120111120111111111111111111111111111111	1:2400-2377	
HAMJ 281	29	ttatttetteeatatggaagtacetatettggaaaaagaa	SEQ ID	NdeI
1171111 201	2)	Hattivitovataiggaagtuvottivitggaaaatgaa	1:2398-2421	
HAMJ 300	30	ttatttetteeatatggtgeetatgeactaaaceage	SEQ ID 1 :62-	NdeI
HAMI 300	30	liaiticitecatatggtgcctatgcactaaaccage	82 B 1 .02-	Taca
	<u></u>		UZ	

HAMJ 313	21	ataagaatgcggccgcttccacaatataagtcgattgatt	SEQ ID	NotI
HAMI 313	31	ataagaatgeggeegetteeacaatataagtegattgatt	1 :2400-2377	
OCRR 487	32	cagtagatctgtgcttatgaactaggtttgc	SEQ ID 3 :58- 79	BglII
OCRR 488	33	gatcaagettgetgetacetttacttactete	SEQ ID 12 :2577-2556	HindIII
HAMJ 171	34	ctgagatatccgttatcgttcaaacc	SEQ ID 3:1060-1075	EcoRV
HAMJ 251	35	ctgcaagcttttaaaggggaataatacg	SEQ ID 3:1059-1045	HindIII
HAMJ 264	36	cagtagatetgeagaageetteetatetg	SEQ ID 3 :682- 700	BglII
HAMJ 282	37	tegecaagettegttategtteaaaceattggg	SEQ ID 3:1060-1081	HindIII
HAMJ 283	38	ataagaatgeggeegeettaeteteetttaataaageeaat agtt	SEQ ID 3 :2520-2492	NdeI
HAMJ 284	39	catgccatggacattgatagtctcttgaaacagc	SEQ ID 3 :856- 880	NcoI
HAMJ 285	40	cgccaagettettaeteteetttaataaagecaatag	SEQ ID 3:2520-2494	HindIII
HAMJ 286	41	cgacaagettaacatggtcgctagegttacc	SEQ ID 3:2139-2119	HindIII
HAMJ 287	42	cataccatgggcctttatgaggcacctaag	SEQ ID 3 :2014-2034	NcoI
HAMJ 288	43	cgacaagettaagtaaatetteageeteteteag	SEQ ID 3 :2376-2353	HindIII
HAMJ 289	44	gataccatggctagcgaccatgttcaaagaa	SEQ ID 3 :2125-2146	NcoI
HAMJ 290	45	cgccaagettatcatccactaacttgaetttatcac	SEQ ID 3:1533-1508	HindIII
HAMJ 291	46	cataccatggatattcttgccttcttagctccg	SEQ ID 3:1531-1554	NcoI
HAMJ 301	47	catgccatggtgcttatgaactaggtttgc	SEQ ID 3 :59- 79	NcoI
HAMJ 302	48	cgccaagetttagegttaccaaaaccattate	SEQ ID 3 :2128-2107	HindIII
HAMJ 160	49	gtattagatetgtteetatgaaettggtegteacea	SEQ ID 13: 172-196	BglII
HAMJ 186	50	cgcctctagactactgtataggagccgg	SEQ ID 13: 2460-2443	XbaI
HAMJ 292	51	catgccatggaaaacatttcaagccttttacgtg	SEQ ID 11: 754-778	NcoI
HAMJ 293	52	cgacaagcttctgtataggagccggttgactttc	SEQ ID 11: 2457-2434	HindIII
HAMJ 294	53	catgccatggttcgtaaaaataaggcagaccaag	SEQ ID 11: 2038-2062	NcoI

HAMJ 297 54 catgccatggaagcctattggaatgggaag SEQ ID 11: NcoI 622-642

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Lists of truncated <u>BVH-3</u> and <u>BVH-11</u> gene products generated from <u>S. pneumoniae</u> Table 3. SP64

desi BVH- L-BV BVH- BVH- BVH-	no	(encoded amino acids)	ID.NO.	vector
			_	
		BVH-3 w/o ss (21-1039)	55	pSL301
		BVH-3 N'end w/o ss (21-509)	56	pSL301
BVH- BVH- BVH-		BVH-3 N'end (1-509)	57	pET-21(+)
BVH-BVH-		BVH-3 C'end (512-1039)	10	pSL301
BVH-		BVH-3 N'end w/o ss (21-225)	59	pET-32 c(+)
		BVH-11 w/o ss (20-840)	09	pCMV-GH
HAMJ251-OCRR487 BVH-11A		BVH-11 N'end w/o ss (20-353)	61	pET-32 c (+)
HAMJ171-OCRR488 BVH-11B		BVH-11 C'end (354-840)	62	pET-32 a(+)
HAMJ264-OCRR488 BVH-11C		BVH-11 C'end (228-840)	63	pET-32 a(+)
HAMJ278-HAMJ279 NEW1		BVH-3 C'end (472-1039)	64	pET-21b(+)
HAMJ278-HAMJ280 NEW2		BVH-3 C'end (472-800)	65	pET-21b(+)
HAMJ281-HAMJ279 NEW3		BVH-3 C'end (800-1039)	99	pET-21b(+)
HAMJ284-HAMJ285 NEW4		BVH-11 C'end (286-840)	67	pET-21d(+)
HAMJ284-HAMJ286 NEW5		BVH-11 internal (286-713)	68	pET-21d(+)
HAMJ287-HAMJ288 NEW6		BVH-11 internal (672-792)	69	pET-21d(+)
HAMJ285-HAMJ289 NEW7		BVH-11 internal (709-840)	70	pET-21d(+)

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HAMJ284-HAMJ290	NEW8	BVH-11 internal (286-511)	71	pET-21d(+)
HAMJ286-HAMJ291	NEW9	BVH-11 internal (511-713)	72	pET-21d(+)
HAMJ160-HAMJ186	BVH-11-2M	BVH-11-2 w/o ss (20-838)	73	pSL301
HAMJ292-HAMJ293	NEW10	BVH-11-2 C'end (271-838)	74	pET-21d(+)
HAMJ293-HAMJ294	NEW11	BVH-11-2 C'end (699-838)	75	pET-21d(+)
HAMJ282-HAMJ283	BVH-11B	BVH-11 C'end (354-840)	62	pET-21b(+)
HAMJ286-HAMJ297	NEW14	BVH-11-2 internal (227-699)	77	pET-21d(+)
HAMJ300-HAMJ313	NEW15	BVH-3 N'end w/o ss (21-800)	78	pET-21b(+)
HAMJ301-HAMJ302	NEW16	BVH-11 N'end w/o ss (20-709)	79	pET-21d(+)

EXAMPLE 9

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This example describes the isolation of monoclonal antibodies (Mabs) and the use of Mabs to characterize BVH-3, BVH-11 and BVH-11-2 protein epitopes.

Female BALB/c mice (Charles River) were immunized subcutaneously with <u>BVH-3</u>, <u>BVH-11</u> or <u>BVH-11-2</u> gene products from <u>S. pneumoniae</u> strain SP64 in presence of 15 μg of QuilA adjuvant (Cedarlane Laboratories Ltd, Hornby, Canada). One set of mice (fusion experiment 1) were immunized on day 1 and 14 with 25 μ g of affinity purified thioredoxin-His•Tag-BVH-3M fusion protein. A second group of mice (fusion experiment 2) were immunized three times at three-week intervals with 25 μg of affinity purified thioredoxin-His • Tag-BVH-11M. A third group of mice (fusion experiment 3) were immunized on day 1 and day 15 with 25 μg of affinity purified thioredoxin-HisoTag-BVH-11-2M fusion protein. A fourth group of mice (fusion experiment 4) were immunized on day 1 with 25 μg of affinity purified thioredoxin-His•BVH-11B fusion protein and boosted by intravenous injection on day 16 and on day 37 with recombinant BVH-11B in PBS. Three to four days before fusion, mice were injected intravenously with 25 $\mu \mathrm{g}$ of the respective antigen suspended in PBS alone. Hybridomas were produced by fusion of spleen cells with nonsecreting SP2/0 myeloma cells as previously described by J. Hamel et al. [J. Med. Microbiol., 23, pp163-170 (1987)]. Culture supernatants of hybridomas were initially screened by enzyme-linked-immunoassay according to the procedure described by Hamel et al. (Supra) using plates coated with

preparations of purified recombinant proteins or suspensions of heat-killed <u>S. pneumoniae</u> cells. Positive hybridomas selected on the basis of ELISA reactivity with a variety of antigens were then cloned by limiting dilutions, expanded and frozen.

Hybridomas were tested by ELISA or Western immunoblotting against BVH-3 and BVH-11 gene products in order to characterize the epitopes recognized by the Mabs. BVH-3 and BVH-11 shared common epitopes with 6 Mabs (H3-1-F9, H3-1-D4, H3-1-H12, H11-1-E7, H11-1-H10 and H11-1.1-G11) showing reactivities with both proteins (Table 4). BVH-11 and BVH-11-2 molecules from <u>S. pneumoniae</u> SP64 shared common epitopes not present on BVH-3 with Mabs (3A1, 13C11, 10H10, 1D8, 10G9, 10A2, 3E8, 10D7, 2H7 and 6H7) reactive with both, BVH-11 and BVH-11-2, recombinant proteins (Table 5).

Table 4. Reactivity of BVH-3-immunoreactive Mabs with a panel of $\underline{\text{BVH-3}}$ and $\underline{\text{BVH-11}}$ gene products

	a.Immunoreactivity with						
MAbs	BVH-3M	BVH-3A	BVH-3B	BVH-3C	NEW2	NEW3	BVH-11M
!	21-1039	21-509	512-1039	21-225	472-800	800-1039	20-840
H3-1-F9	+	+	_	+	-	_	+
H3-1-D4	+	+	_	+	-	-	+
H3-1-H12	+	+	-	+	_	-	+
H3-2-G2	+	+	_	_	_	_	-
H3-3-A1	+	+	_	-	_		-
H3-4-D3	+	-	+	_	-	+	-
H11-1-E7	+	+	-	+	_	_	+
H11-1-	+	+	_	+	_	-	+

H10							
H11-	+	+	_	+	+	_	+
1.1-G11							

Table 5. Reactivity of Mabs raised against BVH-11-2 protein from <u>S. pneumoniae</u> strain SP64 with a panel of <u>BVH-</u>

5 <u>11</u> gene products

	b.Immunoreactivity with							
Mabsa	c.BVH-11 products			d.BVH-11-2 products				
	BVH-11M	NEW8	NEW9	BVH-11B	BVH-11-2	NEW10	NEW11	NEW14
	20-840	286-511	511-713	354-840	20-838	271-838	699-838	227-699
3A1	+	+	-	+	+	+	_	+
13C1	+	+	+	+	+	+	-	+
10H10	+	+	+	+	+	+		+
1D8	+	+	-	+	+	+	-	+
10G9	+	_	_	+	+	+	-	+
10A2	+	_	-	+	+	+	-	+
3E8	+	-	-	+	+	+	-	+
10D7	+	-	_	+	+	+	-	+
2H7	+	-	-	_	+	_	<u> </u>	-
6H7	+	-	-	-	+	_	-	-
3A4	-	-	-	_	+	+	+	-
14H6	-	_	-	-	+	+	+	_
7G2	-	-	_	_	+	+	-	+
13H10	-	_	-	_	+		_	+
7E8	_	_	-	-	+	_	_	_
7H6	-	_		_	+		-	<u> </u>

^a Mabs listed in this table were not reactive with recombinant BVH-3 molecule

The results obtained from the immunoreactivity studies of the Mabs (Table 4 and Table 5) are in agreement with the protein sequences derived from the respective gene sequences. Indeed the Mabs cross-reactive with BVH-3 and BVH-11 molecules recognized BVH-3C protein corresponding to the conserved region, and BVH-11 and BVH-11-2 specific Mabs

were reactive with epitopes located on variable parts of these molecules. BVH-3 and BVH-11, and BVH-11 and BVH-11-2 can be distinguished by their reactivity with Mabs.

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EXAMPLE 10

This example illustrates the simultaneous expression of 10 <u>BVH-3</u> and <u>BVH-11</u> gene products by <u>S. pneumoniae</u>.

A standard Western blot technique was used to investigate whether <u>BVH-3</u> and <u>BVH-11</u> genes were expressed in <u>S.</u> pneumoniae. S. pneumoniae strain SP64 and SP63 were grown overnight at 37°C in 5% CO_2 on chocolate agar plates, bacteria were suspended in PBS and heat-killed at 56°C for 20 min. For the preparation of antigens, suspensions of \underline{S} . pneumoniae were treated with sample buffer containing SDS and 2-mercaptoethanol for 5 min at 100°C. Pneumococcal protein antigens were resolved by SDS-PAGE electrophoresis according to the method of Laemmli [Nature, 227, pp. 680-685 (1970)]. After SDS-PAGE, the proteins were transferred electrophoretically from the gel to nitrocellulose paper by the method of Towbin [Proc. Natl. Acad. Sci. USA, 76, pp. 4350-4354 (1979)] and probed with mouse antiserum or The detection of antigens reactive monoclonal antibodies. with the antibodies was performed by indirect enzymeimmunoassay using conjugated-anti-mouse immunoglobulins and a colour substrate. When antiserum raised to recombinant BVH-3 was tested against <u>S. pneumoniae</u> SP64 antigens, two reactive bands having apparent molecular masses of 127 kDa and 99 kDa were detected. Bands having the same apparent

molecular masses were also detected when Mabs H3-1-F9, H3-1-D4, H3-1-H12, H11-1-E7, H11-1-H10 and H11-1.1-G11 were used individually as immunological probes. In contrast, Mabs specific for the BVH-3 molecule detected the 127 kDa 5 band only and Mabs specific for BVH-11 detected the 99 kDa band only thus confirming the identity of the 127 and 99 kDa bands as BVH-3 and BVH-11, respectively. These studies provide evidence that BVH-3 and BVH-11 proteins are simultaneously present on S. pneumoniae. Moreover, the results are consistent with our previous observations that 10 BVH-3 and BVH-11 possess epitopes that are common to both proteins and epitopes that are exclusive to either protein.

In S. pneumoniae SP64, mature BVH-3, BVH-11 and BVH-11-2 are proteins of 1019, 821 and 819 amino acids with predicted molecular mass of 112.5 kDa, 92.4 kDa, and 91.7 kDa, respectively. Although there is a discrepancy between the molecular mass predicted from the sequence and the molecular mass calculated on SDS-PAGE, BVH-3 can be distinguished from BVH-11 by its higher molecular mass. 20 Moreover, BVH-3 molecules from <u>S. pneumoniae</u> strain SP63 have an apparent molecular mass of 112 kDa in SDS-PAGE compared to 127 kDa for BVH-3 of SP64 strain. This data is consistent with the deletion of a stretch of 177 amino acid residues in BVH-3 of <u>S. pneumoniae</u> strain SP63. 25

EXAMPLE 11

This example describes the protection conferred in 30 experimental infection of mice vaccinated with recombinant BVH-3 or BVH-11 gene products.

Groups of 7 or 8 female BALB/c mice (Charles River) were immunized subcutaneously three times at three-week intervals with either affinity purified thioredoxin-

His•Tag-BVH-3M fusion protein, affinity purified thioredoxin-His•Tag-BVH-11M fusion protein or, as control, with QuilA adjuvant alone in PBS. Twelve to 14 days following the third immunization, the mice were challenged intravenously with <u>S. pneumoniae</u> WU2 strain or intranasally with P4241 strain. Samples of the <u>S. pneumoniae</u> challenge inoculum were plated on chocolate agar plates to determine the CFU and to verify the challenge dose. The challenge dose was approximately 10⁶ CFU. Deaths were recorded for a period of 14 days and on day 14 post-challenge, the surviving mice were sacrificed and blood samples tested for the presence of <u>S. pneumoniae</u> organisms. The survival data are shown in Tables 6 and 7.

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Table 6. Protection mediated by recombinant BVH-3M and BVH-11M proteins in experimental infection with virulent \underline{S} . pneumoniae WU2

Experiment	Immunogen	Alive : dead ^a	Median days alive
1	BVH-3M	8 : 0	>14
	none	0 : 8	1
2	BVH-11M	8 : 0	>14
	none	0 : 8	1

²⁵ a The number of mice alive : the number of mice dead on day 14 post-challenge.

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Table 7. Protection mediated by recombinant BVH-3M and BVH-11M proteins in experimental pneumonia with virulent \underline{S} . pneumoniae P4241

Experiment	Immunogen	Alive : deada	Median day alive
1	BVH-3M	6 : 1	>14
	none	1 : 7	4.5
2	BVH-3M	8:0	>14
	BVH-11M	8 : 0	>14
	none	0:8	4

The number of mice alive: the number of mice dead on day 14 post-challenge.

All mice immunized with recombinant BVH-3M or BVH-11M protein survived to infection with WU2 while none of the control mice given adjuvant alone survived. All except one mice immunized with recombinant BVH-3M or BVH-11M protein survived to infection with P4241 while only one control mice given adjuvant alone survived. All hemocultures from surviving mice were negative at day 14 post-challenge.

These results clearly indicate that both, BVH-3M and BVH-11M, elicit protective anti-pneumococcal immune responses in mice. The fact that these proteins are highly conserved among <u>S. pneumoniae</u> isolates emphasize the potential of BVH-3 and BVH-11 as universal vaccine candidates. Indeed, the BVH-3 and BVH-11 proteins from serogroup 6 <u>S. pneumoniae</u> strain SP64 elicited protection against pneumococcal infections with strains of different capsular serotypes.

25 Ideally, a vaccine that could protect against pneumococcal disease, could protect against meningitis, otitis media,

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bacteremia and pneumonia. BVH-3 and BVH-11 were protective against lethal systemic- and pneumonia-infection models thus suggesting that, in humans, BVH-3- and BVH11-protein-based vaccines could reduce the incidence of a wide spectrum of disease caused by virtually all <u>S. pneumoniae</u> independently of the capsular serotype.

Data from Tables 6 and 7 clearly demonstrate that BVH-3 and BVH-11 were, both, protection-eliciting molecules of <u>S.</u>

<u>pneumoniae</u>. It was not known, however, whether protection can be mediated by specific sequences that were not shared on BVH-3 and BVH-11 molecules. Groups of female BALB/c mice (Charles River) were immunized subcutaneously three times at three-week intervals with either affinity purified thioredoxin-His•Tag-BVH-3AD, -BVH-3B or -BVH-3C fusion protein in presence of 15 μ g of QuilA adjuvant (Cedarlane Laboratories Ltd, Hornby, Canada). Control mice were immunized with QuilA adjuvant alone in PBS or affinity purified thioredoxin-His•Tag or thioredoxin-His•Tag-fusion protein (His-Thio) in presence of QuilA.

To determine the protective ability of a set of truncated proteins, termed NEW4, NEW5, NEW6, NEW7, NEW8, NEW9, NEW10, NEW11, NEW14 and BVH-11B, groups of female BALB/c mice

25 (Charles River) were immunized subcutaneously two times at three-week intervals with 25 μg of either affinity purified His•Tag-fusion protein in presence of 15 μg of QuilA adjuvant. Ten to 14 days following the last immunization, the mice were challenged with virulent S. pneumoniae. Our results indicate that, BVH-3B, a truncated BVH-3 molecule consisting of amino acids 512-1039, elicited protection

against the mouse-virulent strains WU2 and P4241.
Similarly, BVH-11B, NEW4 and NEW5 molecules, three
truncated BVH-11 molecules consisting of amino acids 354840, amino acids 286-840 and amino acids 286-713,

- respectively, elicited protection against experiment intravenous challenge with WU2 and intranasal challenge with P4241. Moreover, vaccination with NEW10 and NEW14, consisting of amino acids 272-838 and amino acids 227-699 from BVH-11-2 molecule also resulted in protection against
- 10 death with the pneumococcal strains. These results indicate that the region comprising 428 amino acids extending from amino acids 286-713 and amino acids 272-699 on <u>S. pneumoniae</u> SP64 BVH-11 and BVH-11-2 protein sequences, respectively, contains protective epitopes.
- 15 This region is highly conserved with a global 91% identity and 94% homology among thirteen BVH-11 protein sequences.

Table 8. Evaluation of protection elicited by vaccination of mice with $\underline{BVH-3}$ and $\underline{BVH-11}$ gene products

		Challenge with WU2		Challenge with P4241	
Experiment	Immunogen	Alive: deada	Median day	Alive : dead	Median day
			alive		alive
1 ^b	None	0 : 8	1.5	1 : 7	4.5
	NEW4	8 : 0	>14	8:0	>14
	NEW5	8 : 0	>14	8 : 0	>14
	NEW7	0 : 8	2	0:8	5
	BVH-11M	8 : 0	>14	8 : 0	>14
2 ^b	None	0:8	1	0:8	4
	NEW5	8 : 0	>14	8 : 0	>14
	NEW8	0:8	1.5	0:8	5.5
	NEW9	3 : 5	3.5	2 : 6	7
	BVH-11M	8 : 0	>14	8 : 0	>14
3 _p	None	0:8	1	0:8	4
	NEW6	0:8	1	4 : 4	10.5 ^c
	NEW10	8 : 0	>14	8 : 0	>14
	NEW11	0 : 8	1.5	1 : 7	6
	BVH-11M	8 : 0	>14	8 : 0	>14
4 ^b	None	0:8	2	0:8	4
	BVH-11B	7 : 1	>14	8 : 0	>14
	NEW14	8 : 0	>14	8 : 0	>14
5	His-Thio	0:8	2		
	BVH-3AD	1 : 7	2.5		
	BVH-3B	5 : 3	>14		
6	His-Thio	0:8	1		
	BVH-3C	0:8	1		

The number of mice alive : the number of mice dead on day

14 post-challenge.

 $^{^{\}rm b}$ The WU2 challenge dose was $10^{\rm 5}$ CFU.

^c Mice living longer than 14 days were assigned a survival time of 14 days for the determination of median values.

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EXAMPLE 12

This example described the cloning and expression of a chimeric gene encoding for a chimeric polypeptide corresponding to the carboxy-terminal region of BVH-3 in fusion at the C' end to the carboxy-terminal region of BVH-11 and the additive protection observed after vaccination with a chimeric polypeptide.

It is clear from the studies described above that BVH-3 and 15 BVH-11 are serologically distinct molecules simultaneously present on S. pneumoniae. The results of immunological studies of mice indicate that both proteins are good vaccine candidates. These proteins have the potential to 20 provide protection against all pneumococci, regardless of serotype. Even though the two proteins share epitopes and sequences, they have different characteristics and may serve different biological functions. Thus, immunization against the two proteins may provide a higher level of protection than that imparted by each individually. examine this, several avenues where full-length or truncated BVH-3 and BVH-11 are administered in combination or in conjugation can be explored. Here we describe the genetic engineering of a BVH-3-BVH-11 fusion gene and protein, termed NEW12 (SEQ ID NO:76 and SEQ ID NO:58, respectively), and the potential use of NEW12 protein as a vaccine.

BVH-3 and BVH-11 gene fragments corresponding to the 3'end of the genes were amplified by PCR using pairs of oligonucleotides engineered to amplify fragments spanning nucleotides 1414 to 3117 (SEQ ID NO: 1) and nucleotides 1060 to 2520 (SEQ ID NO: 3) from S. pneumoniae strain SP64 BVH-3 and BVH-11 genes, respectively. The primers used, HAMJ278 and HAMJ279; HAMJ282 and HAMJ283 had a restriction endonuclease site at the 5' end, thereby allowing directional in-frame cloning of the amplified product into the digested pET21b(+) plasmid vector (Table 2). PCRamplified products were digested with restriction endonucleases and ligated to linearized plasmid pET21b(+) vector digested likewise. The resultant plasmid constructs were confirmed by nucleotide sequence analysis. recombinant pET21b(+) plasmid containing the NdeI-HindIII BVH-3 PCR product was linearized by digestion with the restriction enzymes HindIII and NotI for the in-frame cloning of the HindIII-NotI DNA fragment obtained from the recombinant pET21(+) vector containing the BVH-11 gene fragment. Clones were first stabilized in E. coli DH5 α before introduction into E. coli BL21(λ DE3) for expression of a chimeric pneumococcal protein molecule. The recombinant chimeric polypeptide, termed NEW 12, was expressed as C-terminal fusion with an His-tag. expressed recombinant NEW 12 protein was purified from supernatant fractions obtained from centrifugation of sonicated IPTG-induced E. coli cultures using a His-Bind metal chelation resin (QIAgen, Chatsworth, CA).

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According to the same procedure described above, it is possible to construct other chimeric polypeptides, as a result of a simultaneous expression of New 1 and New 4, New 1 and New 5, New 1 and New 10, or New 1 and New 14. The construction can be with New 1 upstream or downstream of New 4, New 5, New 10, BVH-11B or New 14. It is also possible to construct other chimeric polypeptides as a result of a simultaneous expression of more than two fragments of either genes of BVH-3, BVH-11 or BVH-11-2.

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Groups of 8 female BALB/c mice (Charles River) were immunized subcutaneously two times at three-week intervals with 25 μg of either affinity purified His•Tag-fusion NEW1, BVH-11B or NEW12 protein in presence of 15 μg of QuilA adjuvant. Ten to 14 days following the last immunization, the mice were challenged with virulent <u>S. pneumoniae</u>. demonstrated before, NEW1 and BVH-11B molecules comprising amino acids 472 to 1039 from BVH-3 protein and amino acids 354-840 from BVH-11 protein, respectively, correspond to portions of the proteins capable of eliciting a protective immune response. To determine if a chimeric polypeptide would significantly improve the protection compared with those seen for the individual counterparts, the challenge dose was adjusted in a manner that protection was not expected with NEW1 and BVH-11B molecules. Interestingly, the chimeric NEW12 protein, elicited protection against the Seven out of 8 mice mouse-virulent strains WU2 and P4241. immunized with NEW12 were still alive 10 days after the challenge while 28 out of 32 mice immunized with NEW1, BVH-11B, BVH-3M or adjuvant alone were dead by five days postchallenge. Thus, vaccination of mice with NEW12 provided the highest degree of protection against WU2 challenge.

These results indicate that immunization with a chimeric polypeptide and possibly a combination of BVH-3 and BVH-11 gene products can provide additional protection to that obtained by administration of BVH-3 or BVH-11 antigens alone.

Table 9. Evaluation of protection elicited by vaccination of mice with the chimeric NEW12 molecule

	Challenge with WU2		Challenge with P4241	
Immunogen	Alive : deada	Median day	Alive: dead	Median day
		alive		alive
None	0:8	1	0:8	5
NEW1	2 : 6	2	1 : 7	8
BVH-11B	1 : 7	3.5	8 : 0	>14
NEW12	6 : 2	>14	7 : 1	>14
BVH-3M	1 : 7	3	8 : 1	>14

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EXAMPLE 13

This example illustrates the identification of additional

BVH-3 and BVH-11 related sequences in Streptococcus species other than S. pneumoniae.

It was previously shown that BVH-3, BVH-11 and BVH-11-2 are a family of related proteins sharing common sequences.

20 Homology searches were performed with the nucleotide sequence from the conserved region of these genes and compared with GenBank and EMBL sequences using FASTA. The most significant homology was observed with a 2.469-kb gene coding for a calculated 92-kDa protein (SEQ ID NO: 81) of

unknown function in S. agalactiae also called group B streptococcus or GBS. The gene was designated BVH-71. protein demonstrating 99.2% identity and 99.5% similarity with that of GBS was also identified in S. pyogenes also called group A streptococcus or GAS (SEQ ID NO: 83). The 5' region of the BVH-71 sequences (SEQ ID NO: 80 and SEQ ID NO: 82), spanning nucleotides 1 to 717, demonstrated 58 and 60% identity with the conserved regions of BVH-3 (nucleotides 1 to 675) and <u>BVH-11</u> (nucleotides 1 to 684) 10 genes respectively. The first 239 amino acids of the translated sequences of the GBS and GAS BVH-71 open reading frames are 51 and 54% identical to the first 225 and 228 amino acids of BVH-3 and BVH-11, respectively. In addition to structural similarities, streptococcal BVH-3, BVH-11 and 15 BVH-71 proteins also share antigenic epitopes. A 97-kDa band was revealed on Western blots of GAS or GBS whole cells, using Mab H11-1.1-G11 reactive with the BVH-3 and BVH-11 conserved regions. Similarly, GAS and GBS recombinant BVH-71 proteins were detected in Western 20 immunoblot analysis. These results indicate that BVH-71, BVH-3 and BVH-11 proteins might share similar functions. Our results also suggest that BVH-71 proteins can be used as protein vaccine components of anti-streptococcus. In a further embodiment

BVH-71 proteins can be used as protein vaccine components

of anti-GAS or anti-GBS vaccines.

What is claimed is:

- 1. An isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide having a sequence chosen from: SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.
- 2. A polynucleotide according to claim 1, wherein said polynucleotide encodes a polypeptide having at least 95% identity to the second polypeptide.
- 3. An isolated polynucleotide encoding a polypeptide capable of generating antibodies having binding specificity for a polypeptide having a sequence chosen from: SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.
- 4. An isolated polynucleotide that is complementary to the polynucleotide of claim 1.
- 5. An isolated polynucleotide that is complementary to the polynucleotide of claim 3.
- 6. The polynucleotide of claim 1, wherein said polynucleotide is DNA.
- 7. The polynucleotide of claim 3, wherein said polynucleotide is DNA.
- 8. The polynucleotide of claim 1, wherein said polynucleotide is RNA.

- 9. The polynucleotide of claim 3, wherein said polynucleotide is RNA.
- 10. A vector comprising the polynucleotide of claim 1, wherein said DNA is operably linked to an expression control region.
- 11. A vector comprising the polynucleotide of claim 3, wherein said DNA is operably linked to an expression control region.
- 12. A host cell transfected with the vector of claim 10.
- 13. A host cell transfected with the vector of claim 11.
- 14. A process for producing a polypeptide comprising culturing a host cell according to claim 12 under conditions suitable for expression of said polypeptide.
- 15. A process for producing a polypeptide comprising culturing a host cell according to claim 13 under condition suitable for expression of said polypeptide.
- 16. An isolated polypeptide having at least 70% identity to a second polypeptide having an amino acid sequence chosen from: SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.
- 17. An isolated polypeptide capable of generating antibodies having binding specificity for a second polypeptide

having a sequence chosen from: SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

- 18. An isolated polypeptide having an amino acid sequence chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.
- 19. An isolated polypeptide according to claim 18, wherein the N-terminal Met residue is deleted.
- 20. An isolated polypeptide according to claim 18, wherein the secretory amino acid sequence is deleted.
- 21. A chimeric polypeptide comprising two or more polypeptides chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof; provided that the polypeptides or fragments, analogs or derivatives thereof are linked as to form a chimeric polypeptide.
- 22. A chimeric polypeptide comprising two or more polypeptides chosen from SEQ ID NOs:10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives thereof; provided that the polypeptides or fragments, analogs or derivatives thereof are linked as to form a chimeric polypeptide.

m is 0 or 1,

n is 0 or 1,

A is chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof;

B is chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof;

C is chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof; and

D is chosen from SEQ ID NOs: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

24. A chimeric polypeptide of formula (I):

 $A-(B)_m-(C)_n-D$ (I)

Wherein;

m is 0 or 1,

n is 0 or 1,

A is chosen from SEQ ID NOs :10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives thereof;

B is chosen from SEQ ID NOs :10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77, or fragments, analogs or derivatives thereof;

C is chosen from SEQ ID NOs :10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives thereof; and

D is chosen from SEQ ID NOs :10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives thereof.

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- 25. A vaccine composition comprising a polypeptide according to any one of claims 16 to 24 and a pharmaceutically acceptable carrier, diluent or adjuvant.
- 26. A method for therapeutic or prophylactic treatment of meningitis, otitis media, bacteremia or pneumonia infection in an individual susceptible to meningitis, otitis media, bacteremia or pneumonia infection comprising administering to said individual a therapeutic or prophylactic amount of a composition according to claim 25.
- 27. A method for therapeutic or prophylactic treatment of streptococcal bacterial infection in an individual susceptible to streptococcal infection comprising administering to said individual a therapeutic or prophylactic amount of a composition according to claim 25.
- 28. A method according to claim 26, wherein said individual is a mammal.
- 29. A method according to claim 27, wherein said individual is a human.
- 30. A method according to claim 22, wherein said bacterial infection is <u>S.pneumoniae</u>, group A streptococcus (pyogenes), group B streptococcus (GBS or agalactiae), dysgalactiae, uberis, nocardia or Staphylococcus aureus.

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31. A method according to claim 26, wherein said bacterial infection is <u>S.pneumoniae</u>.

ABSTRACT

f q f t

Streptococcus proteins and polynucleotides encoding them are disclosed. Said proteins are antigenic and therefore useful vaccine components for the prophylaxis or therapy of streptococcus infection in animals. Also disclosed are recombinant methods of producing the protein antigens as well as diagnostic assays for detecting streptococcus bacterial infection.

ATGAAATTTA	GTAAAAAATA	TATAGCAGCT	GGATCAGCTG	TTATCGTATC	CTTGAGTCTA	60
TGTGCCTATG	CACTAAACCA	GCATCGTTCG	CAGGAAAATA	AGGACAATAA	TCGTGTCTCT	120
TATGTGGATG	GCAGCCAGTC	AAGTCAGAAA	AGTGAAAACT	TGACACCAGA	CCAGGTTAGC	180
CAGAAAGAAG	GAATTCAGGC	TGAGCAAATT	GTAATCAAAA	TTACAGATCA	GGGCTATGTA	240
		TCATTACTAT				300
		GGATCCAAAC				360
GAAGTCAAGG	GTGGTTATAT	CATCAAGGTC	GATGGAAAAT	ATTATGTCTA	CCTGAAAGAT	420
CCACCTCATG	CTGATAATGT	TCGAACTAAA	GATGAAATCA	ATCGTCAAAA	ACAAGAACAT	480
CTCAAACATA	ATGAGAAGGT	TAACTCTAAT	GTTGCTGTAG	CAAGGTCTCA	GGGACGATAT	540
ACCACADATC	ATGGTTATGT	CTTTAATCCA	GCTGATATTA	TCGAAGATAC	GGGTAATGCT	600
TATATCCTTC	CTCATGGAGG	TCACTATCAC	TACATTCCCA	AAAGCGATTT	ATCTGCTAGT	660
CAATTACCAC	CACCTAAACC	ACATCTGGCT	GGAAAAAATA	TGCAACCGAG	TCAGTTAAGC	720
TARTIAGCAG	CAGCTAGAGC	CAATAACACG	CAATCTGTAG	CAAAAGGATC	AACTAGCAAG	780
TATICITCAA	AATCTCAAAA	TCTCCAGAGT	CTTTTGAAGG	AACTCTATGA	TTCACCTAGC	840
CCAGCAAAIA	AAICIGAAAA	AGATGGCCTG	CTTTTTCACC	CTCCTAAGAT	TATCAGTCGT	900
GCCCAACGTT	ACAGTGAATC	TCCGCATGGC	CACCATTACC	A CTTTTATTCC	TTACAGCAAG	960
ACACCAAATG	GAGTTGCGAT	CAMMOCCACACA	ATCCTCCCTA	TCACTCGAAC	TGGTTCTACA	1020
CTTTCTGCTT	TAGAAGAAAA	TAATGAAGTA	CTCTCTACTC	TAGGCAGTCT	TTCAAGCAAT	1080
GTTTCTACAA	ATGCAAAACC	TAAGGAGCTC	TCTTCACCAT	CTCATCCTTA	ጥልጥጥጥጥልጥ	1140
CCTTCTTCTT	TAACGACAAG	AACGGCTACA	COMMANAGERI	TANCACATIC	TCTTTTTTT	1200
CCAAAAGATA	TCGTTGAAGA	AACGGCTACA	GCTIATATIG	THACACATA	TAGTCTAGCA	1260
CATTACATTC	CAAAATCAAA	TCAAATTGGG	CAACCGACIC	A THUCAAACAA	TAGTCTAGCA	1320
ACACCTTCTC	CATCTCTTCC	AATCAATCCA	GGAACTTCAC	A I GAGAAACA	TORAGRAGAT	1380
GGATACGGAT	TTGATGCTAA	TCGTATTATC	GCTGAAGATG	AATCAGGTTT	1GICAIGAGI	1440
CACGGAGACC	ACAATCATTA	TTTCTTCAAG	AAGGACTTGA	CAGAAGAGCA	AATTAAGGCT	1500
GCGCAAAAAC	ATTTAGAGGA	AGTTAAAACT	AGTCATAATG	GATTAGATTC	TITIGICATO	1560
CATGAACAGG	ATTATCCAGG	TAATGCCAAA	GAAA'I'GAAAG	ATTTAGATAA	AAAAATCGAA	1620
GAAAAAATTG	CTGGCATTAT	GAAACAATAT	GGTGTCAAAC	GTGAAAGTAT	TGTCGTGAAT	
AAAGAAAAAA	ATGCGATTAT	TTATCCGCAT	GGAGATCACC	ATCATGCAGA	TCCGATTGAT	1680
GAACATAAAC	CGGTTGGAAT	TGGTCATTCT	CACAGTÂACT	ATGAACTGTT	TAAACCCGAA	1740
GAAGGAGTTG	CTAAAAAAAGA	AGGGAATAAA	GTTTATACTG	GAGAAGAATT	AACGAATGTT	1800
GTTAATTTGT	TAAAAAATAG	TACGTTTAAT	AATCAAAACT	TTACTCTAGC	CAATGGTCAA	1860
AAACGCGTTT	CTTTTAGTTT	TCCGCCTGAA	TTGGAGAAAA	AATTAGGTAT	CAATATGCTA	1920
GTAAAATTAA	TAACACCAGA	TGGAAAAGTA	TTGGAGAAAG	TATCTGGTAA	AGTATTTGGA	1980
GAAGGAGTAG	GGAATATTGC	AAACTTTGAA	TTAGATCAAC	CTTATTTACC	AGGACAAACA	2040
TTTAAGTATA	CTATCGCTTC	AAAAGATTAT	CCAGAAGTAA	GTTATGATGG	TACATTTACA	2100
GTTCCAACCT	CTTTAGCTTA	CAAAATGGCC	AGTCAAACGA	TTTTCTATCC	TTTCCATGCA	2160
GGGGATACTT	ATTTAAGAGT	GAACCCTCAA	TTTGCAGTGC	CTAAAGGAAC	TGATGCTTTA	2220
GTCAGAGTGT	TTGATGAATT	TCATGGAAAT	GCTTATTTAG	AAAATAACTA	TAAAGTTGGT	2280
GAAATCAAAT	TACCGATTCC	GAAATTAAAC	CAAGGAACAA	CCAGAACGGC	CGGAAATAAA	2340
ATTCCTGTAA	CCTTCATGGC	AAATGCTTAT	TTGGACAATC	AATCGACTTA	TATTGTGGAA	2400
GTACCTATCT	TGGAAAAAGA	AAATCAAACT	GATAAACCAA	GTATTCTACC	ACAATTTAAA	2460
AGGAATAAAG	CACAAGAAAA	CTCAAAACTT	GATGAAAAGG	TAGAAGAACC	AAAGACTAGT	2520
GAGAAGGTAG	AAAAAGAAAA	ACTTTCTGAA	ACTGGGAATA	GTACTAGTAA	TTCAACGTTA	2580
GAAGAAGTTC	CTACAGTGGA	TCCTGTACAA	GAAAAAGTAG	CAAAATTTGC	TGAAAGTTAT	2640
GGGATGAAGC	TAGAAAATGT	CTTGTTTAAT	ATGGACGGAA	CAATTGAATT	ATATTTACCA	2700
TCAGGAGAAG	TCATTAAAAA	GAATATGGCA	GATTTTACAG	GAGAAGCACC	TCAAGGAAAT	2760
GGTGAAAATA	AACCATCTGA	AAATGGAAAA	GTATCTACTG	GAACAGTTGA	GAACCAACCA	2820
асасававта	AACCAGCAGA	TTCTTTACCA	GAGGCACCAA	. ACGAAAAACC	TGTAAAACCA	2880
GAAAACTCAA	CGGATAATGG	AATGTTGAAT	CCAGAAGGGA	ATGTGGGGAG	TGACCCTATG	2940
TTAGATCCAG	CATTAGAGGA	AGCTCCAGCA	GTAGATCCTG	TACAAGAAAA	ATTAGAAAAA	3000
T TAGA T CCAG	GTTACGGATT	AGGCTTAGAT	AGTGTTATAT	TCAATATGGA	TGGAACGATT	3060
TITACAGCIA	TGCCAAGTGG	AGAAGTGATA	AAAAAGAATT	TATCTGATTT	CATAGCGTAA	3120
(SEQ ID NO			GURE 1			
/DEG ID NO						

MKFSKKYIAA	GSAVIVSLSL	CAYALNOHRS	QENKDNNRVS	YVDGSQSSQK		50
SENLTPDQVS	QKEGIQAEQI	VIKITDQGYV	TSHGDHYHYY	NGKVPYDALF		100
SEELLMKDPN	YQLKDADIVN	EVKGGYIIKV	DGKYYVYLKD	AAHADNVRTK		150
DEINROKQEH	VKDNEKVNSN	VAVARSQGRY	TTNDGYVFNP	ADIIEDTGNA		200
YIVPHGGHYH	YIPKSDLSAS	ELAAAKAHLA	GKNMQPSQLS	YSSTASDNNT		250
QSVAKGSTSK	PANKSENLQS	LLKELYDSPS	AQRYSESDGL	VFDPAKIISR		300
TPNGVAIPHG	DHYHFIPYSK	LSALEEKIAR	MVPISGTGST	VSTNAKPNEV		350
VSSLGSLSSN	PSSLTTSKEL	SSASDGYIFN	PKDIVEETAT	AYIVRHGDHF		400
HYIPKSNQIG	QPTLPNNSLA	TPSPSLPINP	GTSHEKHEED	GYGFDANRII		450
AEDESGFVMS	HGDHNHYFFK	KDLTEEQIKA	AQKHLEEVKT	SHNGLDSLSS		500
HEQDYPGNAK	EMKDLDKKIE	EKIAGIMKQY	GVKRESIVVN	KEKNAIIYPH		550
GDHHHADPID	EHKPVGIGHS	HSNYELFKPE	EGVAKKEGNK	VYTGEELTNV		600
VNLLKNSTFN	NQNFTLANGQ	KRVSFSFPPE	LEKKLGINML	VKLITPDGKV		650
LEKVSGKVFG	EGVGNIANFE	LDQPYLPGQT	FKYTIASKDY	PEVSYDGTFT		700
VPTSLAYKMA	SQTIFYPFHA	GDTYLRVNPQ	FAVPKGTDAL	VRVFDEFHGN		750
AYLENNYKVG	EIKLPIPKLN	QGTTRTAGNK	IPVTFMANAY	LDNQSTYIVE		800
VPILEKENQT	DKPSILPQFK	RNKAQENSKL	DEKVEEPKTS	EKVEKEKLSE		850
TGNSTSNSTL	EEVPTVDPVQ	EKVAKFAESY	GMKLENVLFN	MDGTIELYLP		900
SGEVIKKNMA	DFTGEAPQGN	GENKPSENGK	VSTGTVENQP	TENKPADSLP		950
EAPNEKPVKP	ENSTDNGMLN	PEGNVGSDPM	LDPALEEAPA	VDPVQEKLEK		1000
FTASYGLGLD	SVIFNMDGTI	ELRLPSGEVI	KKNLSDFIA	(SEQ ID NO:	2)	1039

FIGURE 2

ATGAAAATCA ATAAAAAATA TCTAGCTGGG TCAGTAGCTA CACTTGTTTT AAGTGTCTGT 60 GCTTATGAAC TAGGTTTGCA TCAAGCTCAA ACTGTAAAAG AAAATAATCG TGTTTCCTAT 120 ATAGATGGAA AACAAGCGAC GCAAAAAACG GAGAATTTGA CTCCTGATGA GGTTAGCAAG 180 CGTGAAGGAA TCAACGCCGA ACAAATCGTC ATCAAGATTA CGGATCAAGG TTATGTGACC 240 TCTCATGGAG ACCATTATCA TTACTATAAT GGCAAGGTCC CTTATGATGC CATCATCAGT 300 GAAGAGCTCC TCATGAAAGA TCCGAATTAT CAGTTGAAGG ATTCAGACAT TGTCAATGAA 360 ATCAAGGGTG GTTATGTCAT TAAGGTAAAC GGTAAATACT ATGTTTACCT TAAGGATGCA 420 GCTCATGCGG ATAATGTCCG TACAAAAGAA GAAATCAATC GGCAAAAACA AGAACATAGT 480 CAGCATCGTG AAGGAGGGAC TTCAGCAAAC GATGGTGCGG TAGCCTTTGC ACGTTCACAG 540 GGACGCTACA CCACAGATGA TGGTTATATC TTCAATGCAT CTGATATCAT CGAAGATACG 600 660 GGCGATGCCT ATATCGTTCC TCATGGAGAT CATTACCATT ACATTCCTAA GAATGAGTTA TCAGCTAGCG AGTTGGCTGC TGCAGAAGCC TTCCTATCTG GTCGGGAAAA TCTGTCAAAT 720 TTAAGAACCT ATCGCCGACA AAATAGCGAT AACACTCCAA GAACAAACTG GGTACCTTCT 780 840 GTAAGCAATC CAGGAACTAC AAATACTAAC ACAAGCAACA ACAGCAACAC TAACAGTCAA GCAAGTCAAA GTAATGACAT TGATAGTCTC TTGAAACAGC TCTACAAACT GCCTTTGAGT 900 CAACGCCATG TAGAATCTGA TGGCCTTATT TTCGACCCAG CGCAAATCAC AAGTCGAACC 960 GCCAGAGGTG TAGCTGTCCC TCATGGTAAC CATTACCACT TTATCCCTTA TGAACAAATG 1020 TCTGAATTGG AAAAACGAAT TGCTCGTATT ATTCCCCTTC GTTATCGTTC AAACCATTGG 1080 GTACCAGATT CAAGACCAGA AGAACCAAGT CCACAACCGA CTCCAGAACC TAGTCCAAGT 1140 CCGCAACCTG CACCAAATCC TCAACCAGCT CCAAGCAATC CAATTGATGA GAAATTGGTC 1200 AAAGAAGCTG TTCGAAAAGT AGGCGATGGT TATGTCTTTG AGGAGAATGG AGTTTCTCGT 1260 TATATCCCAG CCAAGAATCT TTCAGCAGAA ACAGCAGCAG GCATTGATAG CAAACTGGCC 1320 AAGCAGGAAA GTTTATCTCA TAAGCTAGGA GCTAAGAAAA CTGACCTCCC ATCTAGTGAT 1380 CGAGAATTTT ACAATAAGGC TTATGACTTA CTAGCAAGAA TTCACCAAGA TTTACTTGAT 1440 AATAAAGGTC GACAAGTTGA TTTTGAGGCT TTGGATAACC TGTTGGAACG ACTCAAGGAT 1500 GTCTCAAGTG ATAAAGTCAA GTTAGTGGAT GATATTCTTG CCTTCTTAGC TCCGATTCGT CATCCAGAAC GTTTAGGAAA ACCAAATGCG CAAATTACCT ACACTGATGA TGAGATTCAA 1620 GTAGCCAAGT TGGCAGGCAA GTACACAACA GAAGACGGTT ATATCTTTGA TCCTCGTGAT 1680 ATAACCAGTG ATGAGGGGGA TGCCTATGTA ACTCCACATA TGACCCATAG CCACTGGATT 1740 AAAAAAGATA GTTTGTCTGA AGCTGAGAGA GCGGCAGCCC AGGCTTATGC TAAAGAGAAA 1800 GGTTTGACCC CTCCTTCGAC AGACCATCAG GATTCAGGAA ATACTGAGGC AAAAGGAGCA 1860 GAAGCTATCT ACAACCGCGT GAAAGCAGCT AAGAAGGTGC CACTTGATCG TATGCCTTAC 1920 AATCTTCAAT ATACTGTAGA AGTCAAAAAC GGTAGTTTAA TCATACCTCA TTATGACCAT 1980 TACCATAACA TCAAATTTGA GTGGTTTGAC GAAGGCCTTT ATGAGGCACC TAAGGGGTAT 2040 ACTCTTGAGG ATCTTTTGGC GACTGTCAAG TACTATGTCG AACATCCAAA CGAACGTCCG 2100 CATTCAGATA ATGGTTTTGG TAACGCTAGC GACCATGTTC AAAGAAACAA AAATGGTCAA 2160 GCTGATACCA ATCAAACGGA AAAACCAAGC GAGGAGAAAC CTCAGACAGA AAAACCTGAG 2220 GAAGAAACCC CTCGAGAAGA GAAACCACAA AGCGAGAAAC CAGAGTCTCC AAAACCAACA 2280 GAGGAACCAG AAGAAGAATC ACCAGAGGAA TCAGAAGAAC CTCAGGTCGA GACTGAAAAG 2340 GTTGAAGAA AACTGAGAGA GGCTGAAGAT TTACTTGGAA AAATCCAGGA TCCAATTATC AAGTCCAATG CCAAAGAGAC TCTCACAGGA TTAAAAAATA ATTTACTATT TGGCACCCAG 2460 GACAACAATA CTATTATGGC AGAAGCTGAA AAACTATTGG CTTTATTAAA GGAGAGTAAG 2520 2523 TAA (SEQ ID NO: 3)

FIGURE 3

MKINKKYLAG	SVATLVLSVC	AYELGLHQAQ	TVKENNRVSY	IDGKQATQKT		50
ENLTPDEVSK	REGINAEQIV	IKITDQGYVT	SHGDHYHYYN	GKVPYDAIIS		100
EELLMKDPNY	QLKDSDIVNE	IKGGYVIKVN	${\tt GKYYVYLKDA}$	AHADNVRTKE		150
EINROKOEHS	QHREGGTSAN	DGAVAFARSQ	GRYTTDDGYI	FNASDIIEDT		200
GDAYIVPHGD	HYHYIPKNEL	SASELAAAEA	FLSGRENLSN	LRTYRRQNSD		250
NTPRTNWVPS	VSNPGTTNTN	TSNNSNTNSQ	ASQSNDIDSL	LKQLYKLPLS		300
QRHVESDGLI	FDPAQITSRT	ARGVAVPHGN	${\tt HYHFIPYEQM}$	SELEKRIARI		350
IPLRYRSNHW	VPDSRPEEPS	PQPTPEPSPS	PQPAPNPQPA	PSNPIDEKLV		400
KEAVRKVGDG	YVFEENGVSR	YIPAKNLSAE	TAAGIDSKLA	KQESLSHKLG		450
AKKTDLPSSD	REFYNKAYDL	LARIHQDLLD	NKGRQVDFEA	LDNLLERLKD		500
VSSDKVKLVD	DILAFLAPIR	HPERLGKPNA	QITYTDDEIQ	VAKLAGKYTT		550
EDGYIFDPRD	ITSDEGDAYV	TPHMTHSHWI	KKDSLSEAER	AAAQAYAKEK		600
GLTPPSTDHQ	DSGNTEAKGA	EAIYNRVKAA	KKVPLDRMPY	NLQYTVEVKN		650
GSLIIPHYDH	YHNIKFEWFD	EGLYEAPKGY	TLEDLLATVK	YYVEHPNERP		700
HSDNGFGNAS	DHVQRNKNGQ	ADTNQTEKPS	EEKPQTEKPE	EETPREEKPQ		750
SEKPESPKPT	EEPEEESPEE	SEEPQVETEK	VEEKLREAED	LLGKIQDPII		800
KSNAKETLTG	LKNNLLFGTO	DNNTIMAEAE	KLLALLKESK	(SEQ ID NO:	4)	840

ATGGAGAATA	TAGACATGTT	TAAATCAAAT	CATGAGCGAA	GAATGCGTTA	TTCCATTCGT	60
AAATTTAGTG	TAGGAGTAGC	TAGCGTAGCT	GTTGCCAGTC	TTTTTATGGG	AAGTGTTGTA	120
CATGCGACAG	AGAAAGAGGG	AAGTACCCAA	GCAGCCACTT	CTTTTAATAG	GGGAAATGGA	180
AGTCAGGCAG	AACAACGTGG	AGAACTCGAT	TTAGAACGAG	ATAAGGCAAT	GAAAGCGGTC	240
AGTGAATATG	TAGGAAAAAT	GGTGAGAGAT	GCCTATGTAA	AATCAGATAG	AAAACGACAT	300
AAAAATACTG	TAGCTCTAGT	TAACCAGTTG	GGAAACATTA	AGAACAGGTA	TTTGAATGAA	360
ATAGTTCATT	CAACCTCAAA	AAGCCAACTA	CAGGAACTGA	TGATGAAGAG	TCAATCAGAA	420
GTAGATGAAG		ATTTGAAAAG		CTTCGTCAAG	TTCAGGATCC	480
TCCACTAAAC	CAGAAACTCC			ATCAAAAACC		540
TCTCCGGATA	CCAAACCAAG	CCCTCAACCA		AACCAAGCGT		600
AATCAGGAAA	AAGAAAAAGC	TAAGCTTGCT	GTAGTAACCT	ACATGAGCAA	GATTTTAGAT	660
GATATACAAA	AACATCATCT	GCAGAAAGAA	AAACATCGTC	AGATTGTTGC	TCTTATTAAG	720
GAGCTTGATG	AGCTTAAAAA	GCAAGCTCTT	TCTGAAATTG	ATAATGTAAA		780
GAAATTGAAA	ATACAGTCCA			ATGCAGTTGT	GACTAAATTC	840
AAAAAAGGCT	TAACTCAGGA	CACACCAAAA	GAACCAGGTA	ACAAAAAACC		900
AAACCAGGTA	TGCAACCAAG	TCCTCAACCA	GAGGTTAAAC	CGCAGCTGGA		960
CCAGAGGTTA	AACCGCAACC	AGAAAAACCA	AAACCAGAGG		GCCGGAAAAA	1020
CCAAAACCAG	AGGTTAAACC	GCAGCCGGAA	AAACCAAAAC	CAGAGGTTAA	ACCGCAGCCG	1080
GAAAAACCAA	AACCAGAGGT	TAAACCGCAG	CCGGAAAAAC		GGTTAAACCG	1140
CAGCCGGAAA		AGAGGTTAAA				1200
AAACCGCAGC				AGCCGGAAAA		1260
GAGGTTAAAC	CGCAGCCGGA	AAAACCAAAA	CCAGAGGTTA	AACCGCAACC		1320
AAACCAGAGG	TTAAACCGCA	ACCAGAAAAA	CCAAAACCAG	ATAATAGCAA		1380
GATGATAAGA	AGCCATCAAC	TACAAATAAT	TTAAGCAAGG			1440
GCTTCAACAA		AACAAATAAA		0.11 1 0 0 0 1 1 1 1	AACTGGATCT	1500
ATTTCAAATC	TAGCACTTGA	AATTGCAGGT	CTTCTTACCT	TGGCGGGGGC	AACCATTCTT	1560
GCTAAGAAAA	GAATGAAATA	G (SEQ ID	NO: 5)			1581

FIGURE 5

MENIDMFKSN	HERRMRYSIR	KFSVGVASVA	VASLFMGSVV	HATEKEGSTQ	50
AATSFNRGNG	SQAEQRGELD	LERDKAMKAV	SEYVGKMVRD	AYVKSDRKRH	100
KNTVALVNQL	GNIKNRYLNE	IVHSTSKSQL	QELMMKSQSE	VDEAVSKFEK	150
DSFSSSSGS	STKPETPQPE	NPEHQKPTTP	SPDTKPSPQP	EGKKPSVPDI	200
NQEKEKAKLA	VVTYMSKILD	DIQKHHLQKE	KHRQIVALIK	ELDELKKQAL	250
SEIDNVNTKV	EIENTVHKIF	ADMDAVVTKF	KKGLTQDTPK	EPGNKKPSAP	300
KPGMQPSPQP	EVKPQLEKPK	PEVKPQPEKP	KPEVKPQPEK	PKPEVKPQPE	350
KPKPEVKPQP	EKPKPEVKPQ	PEKPKPEVKP	QPEKPKPEVK	PQPEKPKPEV	400
KPQPEKPKPE	VKPQPEKPKP	EVKPQPEKPK	PEVKPQPEKP	KPEVKPQPEK	450
PKPDNSKPQA	DDKKPSTTNN	LSKDKQPSNQ	ASTNEKATNK	PKKSLPSTGS	500
ISNLALEIAG	LLTLAGATIL	AKKRMK	(SEQ ID NO	D: 6)	526

ATGAAATTTA	GTAAAAAATA	TATAGCAGCT	GGATCAGCTG	TTATCGTATC	CTTGAGTCTA	60
TGTGCCTATG	CACTAAACCA	GCATCGTTCG	CAGGAAAATA	AGGACAATAA	TCGTGTCTCT	120
TATGTGGATG	GCAGCCAGTC	AAGTCAGAAA	AGTGAAAACT	TGACACCAGA	CCAGGTTAGC	180
CAGAAAGAAG	GAATTCAGGC	TGAGCAAATT	GTAATCAAAA	TTACAGATCA	GGGCTATGTA	240
ACGTCACACG	GTGACCACTA	TCATTACTAT	AATGGGAAAG	TTCCTTATGA	TGCCCTCTTT	300
AGTGAAGAAC	TCTTGATGAA	GGATCCAAAC	TATCAACTTA	AAGACGCTGA	TATTGTCAAT	360
GAAGTCAAGG	GTGGTTATAT	CATCAAGGTC	GATGGAAAAT	ATTATGTCTA	CCTGAAAGAT	420
GCAGCTCATG	CTGATAATGT	TCGAACTAAA	GATGAAATCA	ATCGTCAAAA	ACAAGAACAT	480
GTCAAAGATA	ATGAGAAGGT	TAACTCTAAT	GTTGCTGTAG	CAAGGTCTCA	GGGACGATAT	540
ACGACAAATG	ATGGTTATGT	CTTTAATCCA	GCTGATATTA	TCGAAGATAC	GGGTAATGCT	600
TATATCGTTC	CTCATGGAGG	TCACTATCAC	TACATTCCCA	AAAGCGATTT	ATCTGCTAGT	660
GAATTAGCAG	CAGCTAAAGC	ACATCTGGCT	GGAAAAAATA	TGCAACCGAG	TCAGTTAAGC	720
TATTCTTCAA	CAGCTAGTGA	CAATAACACG	CAATCTGTAG	CAAAAGGATC	AACTAGCAAG	780
CCAGCAAATA	AATCTGAAAA	TCTCCAGAGT	CTTTTGAAGG	AACTCTATGA	TTCACCTAGC	840
GCCCAACGTT	ACAGTGAATC	AGATGGCCTG	GTCTTTGACC	CTGCTAAGAT	TATCAGTCGT	900
ACACCAAATG	GAGTTGCGAT	TCCGCATGGC	GACCATTACC	ACTTTATTCC	TTACAGCAAG	960
CTTTCTGCTT	TAGAAGAAAA	GATTGCCAGA	ATGGTGCCTA	TCAGTGGAAC	TGGTTCTACA	1020
GTTTCTACAA	ATGCAAAACC	TAATGAAGTA	GTGTCTAGTC	TAGGCAGTCT	TTCAAGCAAT	1080
CCTTCTTCTT	TAACGACAAG	TAAGGAGCTC	TCTTCAGCAT	CTGATGGTTA	TATTTTTAAT	1140
CCAAAAGATA	TCGTTGAAGA	AACGGCTACA	GCTTATATTG	TAAGACATGG	TGATCATTTC	1200
CATTACATTC	CAAAATCAAA	TCAAATTGGG	CAACCGACTC	TTCCAAACAA	TAGTCTAGCA	1260
ACACCTTCTC	CATCTCTTCC	AATCAATCCA	GGAACTTCAC	ATGAGAAACA	TGAAGAAGAT	1320
GGATACGGAT	TTGATGCTAA	TCGTATTATC	GCTGAAGATG	AATCAGGTTT	TGTCATGAGT	1380
CACGGAGACC	ACAATCATTA	TTTCTTCAAG	AAGGACTTGA	CAGAAGAGCA	AATTAAGGTG	1440
CGCAAAAACA	TTTAG (SI	EQ ID NO: 7))			1455

FIGURE 7

MKFSKKYIAA	GSAVIVSLSL	CAYALNQHRS	QENKDNNRVS	YVDGSQSSQK	50
SENLTPDQVS	QKEGIQAEQI	VIKITDQGYV	TSHGDHYHYY	${\tt NGKVPYDALF}$	100
SEELLMKDPN	YQLKDADIVN	EVKGGYIIKV	DGKYYVYLKD	AAHADNVRTK	150
DEINRQKQEH	VKDNEKVNSN	VAVARSQGRY	TTNDGYVFNP	ADIIEDTGNA	200
YIVPHGGHYH	YIPKSDLSAS	ELAAAKAHLA	GKNMQPSQLS	YSSTASDNNT	250
QSVAKGSTSK	PANKSENLQS	LLKELYDSPS	AQRYSESDGL	VFDPAKIISR	300
TPNGVAIPHG	DHYHFIPYSK	LSALEEKIAR	MVPISGTGST	VSTNAKPNEV	350
	PSSLTTSKEL				400
HYIPKSNQIG	QPTLPNNSLA	TPSPSLPINP	GTSHEKHEED	GYGFDANRII	450
AEDESGFVMS	HGDHNHYFFK	KDLTEEQIKV	RKNI (SE	Q ID NO: 8)	484

ATGAAAGATT	TAGATAAAAA	AATCGAAGAA	AAAATTGCTG	${\tt GCATTATGAA}$	ACAATATGGT	60
GTCAAACGTG	AAAGTATTGT	CGTGAATAAA	GAAAAAAATG	CGATTATTTA	TCCGCATGGA	120
GATCACCATC	ATGCAGATCC	GATTGATGAA	CATAAACCGG	TTGGAATTGG	TCATTCTCAC	180
AGTAACTATG	AACTGTTTAA	ACCCGAAGAA	GGAGTTGCTA	AAAAAGAAGG	GAATAAAGTT	240
TATACTGGAG	AAGAATTAAC	GAATGTTGTT	${\tt AATTTGTTAA}$	AAAATAGTAC	GTTTAATAAT	300
CAAAACTTTA	CTCTAGCCAA	TGGTCAAAAA	CGCGTTTCTT	TTAGTTTTCC	GCCTGAATTG	360
GAGAAAAAAT	TAGGTATCAA	TATGCTAGTA	AAATTAATAA	CACCAGATGG	AAAAGTATTG	420
GAGAAAGTAT	CTGGTAAAGT	ATTTGGAGAA	GGAGTAGGGA	ATATTGCAAA	CTTTGAATTA	480
GATCAACCTT	ATTTACCAGG	ACAAACATTT	AAGTATACTA	TCGCTTCAAA	AGATTATCCA	540
GAAGTAAGTT	ATGATGGTAC	ATTTACAGTT	CCAACCTCTT	TAGCTTACAA	AATGGCCAGT	600
CAAACGATTT	TCTATCCTTT	CCATGCAGGG	GATACTTATT	TAAGAGTGAA	CCCTCAATTT	660
GCAGTGCCTA	AAGGAACTGA	TGCTTTAGTC	AGAGTGTTTG	ATGAATTTCA	TGGAAATGCT	720
TATTTAGAAA	ATAACTATAA	AGTTGGTGAA	ATCAAATTAC	CGATTCCGAA	ATTAAACCAA	780
GGAACAACCA	GAACGGCCGG	AAATAAAATT	CCTGTAACCT	TCATGGCAAA	TGCTTATTTG	840
GACAATCAAT	CGACTTATAT	TGTGGAAGTA	CCTATCTTGG	AAAAAGAAAA		900
AAACCAAGTA	TTCTACCACA	ATTTAAAAGG	AATAAAGCAC	AAGAAAACTC		960
GAAAAGGTAG	AAGAACCAAA	GACTAGTGAG	AAGGTAGAAA	AAGAAAAACT	TTCTGAAACT	1020
GGGAATAGTA	CTAGTAATTC	AACGTTAGAA	GAAGTTCCTA	CAGTGGATCC	TGTACAAGAA	1080
AAAGTAGCAA	AATTTGCTGA	AAGTTATGGG	ATGAAGCTAG	AAAATGTCTT	GTTTAATATG	1140
GACGGAACAA	TTGAATTATA	TTTACCATCA	GGAGAAGTCA	TTAAAAAGAA	TATGGCAGAT	1200
TTTACAGGAG	AAGCACCTCA	AGGAAATGGT	GAAAATAAAC	CATCTGAAAA	TGGAAAAGTA	1260
TCTACTGGAA	CAGTTGAGAA	CCAACCAACA	GAAAATAAAC	CAGCAGATTC	TTTACCAGAG	1320
GCACCAAACG	AAAAACCTGT	AAAACCAGAA	AACTCAACGG	ATAATGGAAT	GTTGAATCCA	1380
GAAGGGAATG	TGGGGAGTGA	CCCTATGTTA	GATCCAGCAT	TAGAGGAAGC	TCCAGCAGTA	1440
GATCCTGTAC	AAGAAAAATT	AGAAAAATTT	ACAGCTAGTT	ACGGATTAGG	CTTAGATAGT	1500
GTTATATTCA	ATATGGATGG	AACGATTGAA	TTAAGATTGC	CAAGTGGAGA	AGTGATAAAA	1560
AAGAATTTAT	CTGATTTCAT	AGCGTAA	(SEQ ID NO	0: 9)		1587

FIGURE 9

MKDLDKKIEE	KIAGIMKQYG	VKRESIVVNK	EKNAIIYPHG	DHHHADPIDE	50
	SNYELFKPEE				100
ONFTLANGQK	RVSFSFPPEL	EKKLGINMLV	KLITPDGKVL	EKVSGKVFGE	150
	DQPYLPGQTF				200
OTIFYPFHAG	DTYLRVNPQF	AVPKGTDALV	RVFDEFHGNA	YLENNYKVGE	250
	GTTRTAGNKI				300
KPSILPQFKR	NKAQENSKLD	EKVEEPKTSE	KVEKEKLSET	GNSTSNSTLE	350
EVPTVDPVQE	KVAKFAESYG	MKLENVLFNM	DGTIELYLPS	GEVIKKNMAD	400
	ENKPSENGKV				450
NSTDNGMLNP	EGNVGSDPML	DPALEEAPAV	DPVQEKLEKF	TASYGLGLDS	500
	LRLPSGEVIK				528

FIGURE 10

BVH3		1	CAYALNQHRSQENKDNNRVSYVDGSQSSQKSENLTPDQVSQKEGIQAEQIVIKITDQGYV CAYALNQHRSQENKDNNRVSYVDGSQSSQKSENLTPDQVSQKEGIQAEQIVIKITDQGYV	60 60
BVH3	JNR7/87	1	CAYALNQHRSQENKDNNRVSYVDGSQSSQKSENLTPDQVSQKEGIQAEQIVIKITDQGYV	60
BVH3	-	1	CAYALNQHRSQENKDNNRVSYVDGSQSSQKSENLTPDQVSQKEGIQAEQIVIKITDQGYV	60
	P4241	1	CAYALNQHRSQENKDNNRVSYVDGSQSSQKSENLTPDQVSQKEGIQAEQIVIKITDQGYV	60
BVH3		1	CAYALNQHRSQENKDNNRVSYVDGSQSSQKSENLTPDQVSQKEGIQAEQIVIKITDQGYV	60
DVID	AOO	_	****************	
BVH3	WU2	61	TSHGDHYHYYNGKVPYDALFSEELLMKDPNYQLKDADIVNEVKGGYIIKVDGKYYVYLKD	120
BVH3	RX1	61	TSHGDHYHYYNGKVPYDALFSEELLMKDPNYQLKDADIVNEVKGGYIIKVDGKYYVYLKD	120
BVH3	JNR7/87	61	TSHGDHYHYYNGKVPYDALFSEELLMKDPNYQLKDADIVNEVKGGYIIKVDGKYYVYLKD	120
BVH3	SP64	61	TSHGDHYHYYNGKVPYDALFSEELLMKDPNYQLKDADIVNEVKGGYIIKVDGKYYVYLKD	120 120
	P4241	61	TSHGDHYHYYNGKVPYDALFSEELLMKDPNYQLKDADIVNEVKGGYIIKVDGKYYVYLKD	120
BVH3	A66	61	TSHGDHYHYYNGKVPYDALFSEELLMKDPNYQLKDADIVNEVKGGYIIKVDGKYYVYLKD ************************************	120
BVH3	WII2	121	AAHADNVRTKDEINRQKQEHVKDNEKVNSNVAVARSQGRYTTNDGYVFNPADIIEDTGNA	180
BVH3		121	AAHADNVRTKDEINROKOEHVKDNEKVNSNVAVARSQGRYTTNDGYVFNPADIIEDTGNA	180
	JNR7/87	121	AAHADNVRTKDEINROKOEHVKDNEKVNSNVAVARSQGRYTTNDGYVFNPADIIEDTGNA	180
BVH3		121	AAHADNVRTKDEINRQKQEHVKDNEKVNSNVAVARSQGRYTTNDGYVFNPADIIEDTGNA	180
BVH3	P4241	121	AAHADNVRTKDEINRQKQEHVKDNEKVNSNVAVARSQGRYTTNDGYVFNPADIIEDTGNA	180
BVH3	A66	121	AAHADNVRTKDEINRQKQEHVKDNEKVNSNVAVARSQGRYTTNDGYVFNPADIIEDTGNA ************************************	180
			YIVPHRGHYHYIPKSDLSASELAAAKAHLAGKNMQPSQLSYSSTASDNNTQSVAKGSTSK	240
BVH3		181	YIVPHRGHYHYIPKSDLSASELAAAKAHLAGKNMQPSQLSYSSTASDNNTQSVAKGSTSK	240
BVH3		181	YIVPHGGHYHYIPKSDLSASELAAKAHLAGKNMQPSQLSYSSTASDNNTQSVAKGSTSK	240
	JNR7/87 SP64	101	YIVPHGGHYHYIPKSDLSASELAAAKAHLAGKNMQPSQLSYSSTASDNNTQSVAKGSTSK	240
	P4241	181	YIVPHRGHYHYIPKSDLSASELAAAKAHLAGKNMQPSQLSYSSTASDNNTQSVAKGSTSK	240
BVH3		181	YTVPHRGHYHYTPKSDLSASELAAAKAHLAGKNMQPSQLSYSSTASDNNTQSVAKGSTSK	240
21110			***** ***************************	
BVH3	WU2	241	PANKSENLQSLLKELYDSPSAQRYSESDGLVFDPAKIISRTPNGVAIPHGDHYHFIPYSK	300
BVH3	RX1	241	PANKSENLQSLLKELYDSPSAQRYSESDGLVFDPAKIISRTPNGVAIPHGDHYHFIPYSK	300 300
	JNR7/87	241	PANKSENLQSLLKELYDSPSAQRYSESDGLVFDPAKIISRTPNGVAIPHGDHYHFIPYSK	300
	SP64	241	PANKSENLQSLLKELYDSPSAQRYSESDGLVFDPAKIISRTPNGVAIPHGDHYHFIPYSK PANKSENLQSLLKELYDSPSAQRYSESDGLVFDPAKIISRTPNGVAIPHGDHYHFIPYSK	300
	P4241	241	PANKSENLQSLLKELYDSPSAQRYSESDGLVFDPAKIISRTPNGVAITHODHYM I SK PANKSENLQSLLKELYDSPSAQRYSESDGLVFDPAKIISRTPNGVAIPHGDHYHFIPYSK	300
BVH3	A66	241	**************************************	
BVH3	WU2	301	LSALEEKIARMVPISGTGSTVSTNAKPNEVVSSLGSLSSNPSSLTTSKELSSASDGYIFN	360
BVH3	RX1	301	LSALEEKIARRVPISGTGSTVSTNAKPNEVVSSLGSLSSNPSSLTTSKELSSASDGYIFN	360
BVH3	JNR7/87	301	LSALEEKIARMVPISGTGSTVSTNAKPNEVVSSLGSLSSNPSSLTTSKELSSASDGYIFN	360 360
BVH3	SP64	301	LSALEEKIARMVPISGTGSTVSTNAKPNEVVSSLGSLSSNPSSLTTSKELSSASDGYIFN	360
	P4241	301	LSALEEKIARMVPISGTGSTVSTNAKPNEVVSSLGSLSSNPSSLTTSKELSSASDGYIFN LSALEEKIARMVPISGTGSTVSTNAKPNEVVSSLGSLSSNPSSLTTSKELSSASDGYIFN	360
BVH3	A66	301	LSALEEKTARMVF15G1G51V51WACENEVV55BG1B5R256B125R256B22224 *********************************	
BVH3	WU2	361	PKDIVEETATAYIVRHGDHFHYIPKSNQIGQPTLPNNSLATPSPSLPINPGTSHEKHEED	420
	RX1	361	PKDIVEETATAYIVRHGDHFHYIPKSNQIGQPTLPNNSLATPSPSLPINPGISHEKHEED	420
	JNR7/87	361	PKDIVEETATAYIVRHGDHFHYIPKSNQIGQPTLPNNSLATPSPSLPINPGTSHEKHEED	420
BVH3	SP64	361	PKDIVEETATAYIVRHGDHFHYIPKSNQIGQPTLPNNSLATPSPSLPINPGTSHEKHEED	420
BVH3	P4241	361	PKDIVEETATAYIVRHGDHFHYIPKSNQIGQPTLPNNSLATPSPSLPINPGTSHEKHEED	420 420
BVH3	A66	361	PKDIVEETATAYIVRHGDHFHYIPKSNQIGQPTLPNNSLATPSPSLPINPGTSHEKHEED	420
נשוזם	WU2	421	GYGFDANRIIAEDESGFVMSHGDHNHYFFKKDLTEEQIKAAQKHLEEVKTSHNGLDSLSS	480
	RX1	421	CYCEDANRITAEDESGFIMSHGNHNHYFFKKDLTEEQIKAAQKHLEEVKTSHNGLDSLSS	480
	JNR7/87	421	GYGFDANRITAEDESGFVMSHGDHNHYFFKKDLTEEQIKAAQKHLEEVKTSHNGLDSLSS	480
	SP64	421	CYCEDANRITAEDESGEVMSHGDHNHYFFKKDLTEEQIKAAQKHLEEVKTSHNGLDSLSS	480
BVH3	P4241	421	GYGFDANRITAEDESGFVMSHGDHNHYFFKKDLTEEQIKAAQKHLEEVKTSHNGLDSLSS	480 480
BVH3	A66	421	GYGFDANRIIAEDESGFVMSHGDHNHYFFKKDLTEEQIKAAQKHLEEVKTSHNGLDSLSS *********************************	480

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481 HEQDYPGNAKEMKDLDKKIEEKIAGIMKQYGVKRESIVVNKEKNAIIYPHGDHHHADPID
BVH3 RX1
              481 HEODYPSNAKEMKDLDKKIEEKIAGIMKQYGVKRESIVVNKEKNAIIYPHGDHHHADPID
BVH3 JNR7/87
              481 HEQDYPGNAKEMKDLDKKIEEKIAGIMKQYGVKRESIVVNKEKNAIIYPHGDHHHADPID
                                                                              540
BVH3 SP64
              481 HEQDYPSNAKEMKDLDKKIEEKIAGIMKQYGVKRESIVVNKEKNAIIYPHGDHHHADPID
                                                                              540
BVH3 P4241
              481 HEQDYPSNAKEMKDLDKKIEEKIAGIMKQYGVKRESIVVNKEKNAIIYPHGDHHHADPID
BVH3 A66
              541 EHKPVGIGHSHSNYELFKPEEGVAKKEGNKVYTGEELTNVVNLLKNSTFNNQNFTLANGQ
                                                                              600
BVH3 WU2
              541 EHKPVGIGHSHSNYELFKPEEGVAKKEGNKVYTGEELTNVVNLLKNSTFNNQNFTLANGQ
                                                                              600
BVH3 RX1
                                                                              600
              541 EHKPVGIGHSHSNYELFKPEEGVAKKEGNKVYTGEELTNVVNLLKNSTFNNQNFTLANGQ
BVH3 JNR7/87
              541 EHKPVGIGHSHSNYELFKPEEGVAKKEGNKVYTGEELTNVVNLLKNSTFNNQNFTLANGQ
                                                                              600
BVH3 SP64
                                                                              600
              541 EHKPVGIGHSHSNYELFKPEEGVAKKEGNKVYTGEELTNVVNLLKNSTFNNQNFTLANGQ
BVH3 P4241
              541 EHKPVGIGHSHSNYELFKPEEGVAKKEGNKVYTGEELTNVVNLLKNSTFNNQNFTLANGQ
                                                                              600
BVH3 A66
              601 KRVSFSFPPELEKKLGINMLVKLITPDGKVLEKVSGKVFGEGVGNIANFELDQPYLPGQT
                                                                              660
BVH3 WU2
              601 KRVSFSFPPELEKKLGINMLVKLITPDGKVLEKVSGKVFGEGVGNIANFELDQPYLPGQT
                                                                              660
BVH3 RX1
              601 KRVSFSFPPELEKKLGINMLVKLITPDGKVLEKVSGKVFGEGVGNIANFELDQPYLPGQT
                                                                              660
BVH3 JNR7/87
              601 KRVSFSFPPELEKKLGINMLVKLITPDGKVLEKVSGKVFGEGVGNIANFELDQPYLPGQT
                                                                              660
BVH3 SP64
              601 KRVSFSFPPELEKKLGINMLVKLITPDGKVLEKVSGKVFGEGVGNIANFELDQPYLPGQT
                                                                              660
BVH3 P4241
              601 KRVSFSFPPELEKKLGINMLVKLITPDGKVLEKVSGKVFGEGVGNIANFELDQPYLPGQT
                                                                              660
BVH3 A66
                   *****************
              661 FKYTIASKDYPEVSYDGTFTVPTSLAYKMASQTIFYPFHAGDTYLRVNPQFAVPKGTDAL
                                                                              720
BVH3 WU2
              661 FKYTIASKDYPEVSYDGTFTVPTSLAYKMASQTIFYPFHAGDTYLRVNPQFAVPKGTDAL
                                                                              720
BVH3 RX1
              661 FKYTIASKDYPEVSYDGTFTVPTSLAYKMASQTIFYPFHAGDTYLRVNPQFAVPKGTDAL
                                                                              720
BVH3 JNR7/87
              661 FKYTIASKDYPEVSYDGTFTVPTSLAYKMASQTIFYPFHAGDTYLRVNPQFAVPKGTDAL
                                                                              720
BVH3 SP64
                                                                              720
              661 FKYTIASKDYPEVSYDGTFTVPTSLAYKMASQTIFYPFHAGDTYLRVNPQFAVPKGTDAL
BVH3 P4241
              661 FKYTIASKDYPEVSYDGTFTVPTSLAYKMASQTIFYPFHAGDTYLRVNPQFAVPKGTDAL
                                                                              720
BVH3 A66
                   ******************
                                                                              780
              721 VRVFDEFHGNAYLENNYKVGEIKLPIPKLNQGTTRTAGNKIPVTFMANAYLDNQSTYIVE
BVH3 WU2
              721 VRVFDEFHGNAYLENNYKVGEIKLPIPKLNQGTTRTAGNKIPVTFMANAYLDNQSTYIVE
                                                                              780
BVH3 RX1
              721 VRVFDEFHGNAYLENNYKVGEIKLPIPKLNQGTTRTAGNKIPVTFMANAYLDNQSTYIVE
BVH3 JNR7/87
                                                                              780
              721 VRVFDEFHGNAYLENNYKVGEIKLPIPKLNQGTTRTAGNKIPVTFMANAYLDNQSTYIVE
BVH3 SP64
              721 VRVFDEFHGNAYLENNYKVGEIKLPIPKLNQGTTRTAGNKIPVTFMANAYLDNQSTYIVE
                                                                              780
BVH3 P4241
                                                                              780
              721 VRVFDEFHGNAYLENNYKVGEIKLPIPKLNQGTTRTAGNKIPVTFMANAYLDNQSTYIVE
BVH3 A66
                   *****************
               781 VPILEKENQTDKPSILPQFKRNKAQENSKFDEKVEEPKTSEKVEKEKLSETGNSTSNSTL
                                                                              840
BVH3 WU2
               781 VPILEKENQTDKPSILPQFKRNKAQENSKLDEKVEEPKTSEKVEKEKLSETGNSTSNSTL
BVH3 RX1
               781 VPILEKENQTDKPSILPQFKRNKAQENLKLDEKVEEPKTSEKVEKEKLSETGNSTSNSTL
                                                                              840
BVH3 JNR7/87
               781 VPILEKENQTDKPSILPQFKRNKAQENSKLDEKVEEPKTSEKVEKEKLSETGNSTSNSTL
                                                                              840
BVH3 SP64
                                                                              840
               781 VPILEKENOTDKPSILPQFKRNKAQENSKFDEKVEEPKTSEKVEKEKLSETGNSTSNSTL
BVH3 P4241
               781 VPILEKENQTDKPSILPQFKRNKAQENSKFDEKVEEPKTSEKVEKEKLSETGNSTSNSTL
                                                                               840
BVH3 A66
                         ***************
               841 BEVPTVDPVQEKVAKFAESYGMKLENVLFNMDGTIBLYLPSGEVIKKNMADFTGEAPQGN
                                                                               900
BVH3 WU2
               841 EEVPTVDPVQEKVAKFAESYGMKLENVLFNMDGTIELYLPSGEVIKKNMADFTGEAPQGN
                                                                               900
BVH3 RX1
               841 EEVPTVDPVQEKVAKFAESYGMKLENVLFNMDGTIELYLPSGEVIKKNMADFTGEAPQGN
                                                                               900
BVH3 JNR7/87
               841 EEVPTVDPVQEKVAKFAESYGMKLENVLFNMDGTIELYLPSGEVIKKNMADFTGEAPQGN
                                                                               900
BVH3 SP64
               841 EEVPTVDPVQEKVAKFAESYGMKLENVLFNMDGTIELYLPSGEVIKKNMADFTGEAPQGN
                                                                               900
BVH3 P4241
               841 EEVPTVDPVQEKVAKFAESYGMKLENVLFNMDGTIELYLPSGEVIKKNMADFTGEAPQGN
                                                                               900
BVH3 A66
                   ***************
               901 GENKPSENGKVSTGTVENQPTENKPADSLPEAPNEKPVKPENSTDNGMLNPEGNVGSDPM
                                                                               960
BVH3 WU2
               901 GENKPSENGKVSTGTVENQPTENKPADSLPEAPNEKPVKPENSTDNGMLNPEGNVGSDPM
BVH3 RX1
               901 GENKPSENGKVSTGTVENQPTENKPADSLPEAPNEKPVKPENSTDNGMLNPEGNVGSDPM
                                                                               960
BVH3 JNR7/87
               901 GENKPSENGKVSTGTVENQPTENKPADSLPEAPNEKPVKPENSTDNGMLNPEGNVGSDPM
BVH3 SP64
               901 GENKPSENGKVSTGTVENQPTENKPADSLPEAPNEKPVKPENSTDNGMLNPEGNVGSDPM
                                                                               960
BVH3 P4241
               901 GENKPSENGKVSTGTVENQPTENKPADSLPEAPNEKPVKPENSTDNGMLNPEGNVGSDPM
                                                                               960
BVH3 A66
                   **************
               961 LDPALEEAPAVDPVQEKLEKFTASYGLGLDSVIFNMDGTIELRLPSGEVIKKNLSDLIA 1019
BVH3 WU2
               961 LDPALEEAPAVDPVQEKLEKFTASYGLGLDSVIFNMDGTIELRLPSGEVIKKNLSDLIA 1019
BVH3 RX1
               961 LDPALEEAPAVDPVQEKLEKFTASYGLGLDSVIFNMDGTIELRLPSGEVIKKNLSDLIA 1019
BVH3 JNR7/87
               961 LDPALEEAPAVDPVQEKLEKFTASYGLGLDSVIFNMDGTIELRLPSGEVIKKNLSDFIA 1019
BVH3 SP64
               961 LDPALEEAPAVDPVQEKLEKFTASYGLGLDSVIFNMDGTIELRLPSGEVIKKNLSDLIA 1019
BVH3 P4241
               961 LDPALEEAPAVDPVQEKLEKFTASYGLGLDSVIFNMDGTIELRLPSGEVIKKNLSDLIA 1019
BVH3 A66
```

FIGURE 11

```
1 CSYELGRHOAGOVKKESNRVSYIDGDQAGQKAENLTPDEVSKREGINAEQIVIKITDQGY
BVH11-2 SP64
                 1 CSYELGRHQAGQVKKESNRVSYIDGDQAGQKAENLTPDEVSKREGINAEQIVIKITDQGY
BVH11-2 JNR7/87
                 1 CSYELGRHQAGQDKKESNRVAYIDGDQAGQKAENLTPDEVSKREGINAEQIVIKITDQGY
                                                                                 60
BVH11-2 P4241
                 1 CSYELGRHQAGQDKKESNRVAYIDGDQAGQKAENLTPDEVSKREGINAEQIVIKITDQGY
BVH11-2 A66
                 1 CSYELGRHQAGQDKKESNRVAYIDGDQAGQKAENLTPDEVSKREGINAEQIVIKITDQGY
                                                                                 60
BVH11-2 WU2
                 1 CSYELGRHOAGOVKKESNRVSYIDGDQAGQKAENLTPDEVSKREGINAEQIVIKITDQGY
BVH11-2 Rx1
                 {\tt 1} {\tt CSYELGRHQAGQDKKESNRVAYIDGDQAGQKAENLTPDEVSKREGINAEQIVIKITDQGY}
                                                                                 60
BVH11 P4241
                 1 CSYELGRHQAGQDKKESNRVAYIDGDQAGQKAENLTPDEVSKREGINAEQIVIKITDQGY
                                                                                 60
BVH11 WU2
                 {\tt 1} {\tt CSYELGRHQAGQDKKESNRVAYIDGDQAGQKAENLTPDEVSKREGINAEQIVIKITDQGY}
                                                                                 60
BVH11 A66
                 1 CSYELGRHOAGOVKKESNRVSYIDGDQAGQKAENLTPDEVSKREGINAEQIVIKITDQGY
                                                                                 60
BVH11 Rx1
                 1 CSYELGRHQAGQDKKESNRVAYIDGDQAGQKAENLTPDEVSKREGINAEQIVIKITDQGY
                                                                                 60
BVH11 JNR7/87
BVH11 SP63
                 1 CSYELGRHQAGQVKKESNRVSYIDGDQAGQKAENLTPDEVSKREGINAEQIVIKITDQGY
                                                                                 60
                 1 CAYELGLHQA-QTVKENNRVSYIDGKQATQKTENLTPDEVSKREGINAEQIVIKITDQGY
BVH11 SP64
                                 ** *** *** ** ** ** ************
                61 VTSHGDHYHYYNGKVPYDAIISEELLMKDPNYQLKDSDIVNEIKGGYVIKVDGKYYVYLK 120
BVH11-2 SP64
                61 VTSHGDHYHYYNGKVPYDAIISEELLMKDPNYQLKDSDIVNEIKGGYVIKVDGKYYVYLK 120
BVH11-2 JNR7/87
                61 VTSHGDHYHYYNGKVPYDAIISEELLMKDPNYQLKDSDIVNEIKGGYVIKVNGKYYVYLK 120
BVH11-2 P4241
                61 VTSHGDHYHYYNGKVPYDAIISEELLMKDPNYQLKDSDIVNEIKGGYVIKVNGKYYVYLK 120
BVH11-2 A66
                61 VTSHGDHYHYYNGKVPYDAIISEELLMKDPNYQLKDSDIVNEIKGGYVIKVNGKYYVYLK 120
BVH11-2 WU2
                61 VTSHGDHYHYYNGKVPYDAIISEELLMKDPNYQLKDSDIVNEIKGGYVIKVDGKYYVYLK 120
BVH11-2 Rx1
                61 VTSHGDHYHYYNGKVPYDAIISEELLMKDPNYQLKDSDIVNEIKGGYVIKVNGKYYVYLK 120
BVH11 P4241
                61 VTSHGDHYHYYNGKVPYDAIISEELLMKDPNYQLKDSDIVNEIKGGYVIKVNGKYYGYLK 120
BVH11 WU2
                61 VTSHGDHYHYYNGKVPYDAIISEELLMKDPNYQLKDSDIVNEIKGGYVIKVNGKYYVYLK 120
BVH11 A66
                61 VTSHGDHYHYYNGKVPYDAIISEELLMKDPNYQLKDSDIVNEIKGGYVIKVDGKYYVYLK 120
BVH11 Rx1
                61 VTSHGDHYHYYNGKVPYDAIISEELLMKDPNYQLKDSDIVNEIKGGYVIKVNGKYYVYLK 120
BVH11 JNR7/87
                61 VTSHGDHYHYYNGKVPYDAIISEELLMKDPNYQLKDSDIVNEIKGGYVIKVDGKYYVYLK 120
BVH11 SP63
                60 VTSHGDHYHYYNGKVPYDAIISEELLMKDPNYQLKDSDIVNEIKGGYVIKVNGKYYVYLK 119
BVH11 SP64
                121 DAAHADNIRTKEEIKRQKQEHSHNHNSRA---DNAVAAARAQGRYTTDDGYIFNASDIIE 177
BVH11-2 SP64
BVH11-2 JNR7/87 121 DAAHADNIRTKEEIKRQKQEHSHNHGGGSN--DQAVVAARAQGRYTTDDGYIFNASDIIE 178
                121 DAAHADNIRTKEEIKRQKQEHSHNHGGGSN--DQAVVAARAQGRYTTDDGYIFNASDIIE 178
BVH11-2 P4241
                121 DAAHADNIRTKEEIKRORQEHSHNHGGGSN--DQAVVAARAQGRYTTDDGYIFNASDIIE 178
BVH11-2 A66
                121 DAAHADNIRTKEEIKROKOEHSHNHGGGSN--DQAVVAARAQGRYTTDDGYIFNASDIIE 178
BVH11-2 WU2
                121 DAAHADNIRTKEEIKRQKQERSHNHNSRA---DNAVAAARAQGRYTTDDGYIFNASDIIE 177
BVH11-2 Rx1
                121 DAAHADNIRTKEEIKROKOEHSHNHGGGSN--DQAVVAARAQGRYTTDDGYIFNASDIIE 178
BVH11 P4241
                121 DAAHADNIRTKEEIKRQKQEHSHNHGGGSN--DQAVVAARAQGRYTTDDGYIFNASDIIE 178
BVH11 WU2
                121 DAAHADNIRTKEEIKRQKQEHSHNHGGGSN--DQAVVAARAQGRYTTDDGYIFNASDIIE 178
BVH11 A66
                121 DAAHADNIRTKEEIKRQKQERSHNHNSRA---DNAVAAARAQGRYTTDDGYIFNASDIIE 177
BVH11 Rx1
               121 DAAHADNIRTKEEIKRQKQERSHNHNSRA---DNAVAAARAQGRYTTDDGYIFNASDIIE 177
BVH11 JNR7/87
                121 DAAHADNIRTKEEIKROKOERSHNHNSRA---DNAVAAARAQGRYTTDDGYIFNASDIIE 177
BVH11 SP63
                120 DAAHADNVRTKEEINRQKQEHSQHREGGTSANDGAVAFARSQGRYTTDDGYIFNASDIIE 179
BVH11 SP64
                    178 DTGDAYIVPHGDHYHYIPKNELSASELAAAEAYWNGKQGSRPSSSSSYNANPVQPRLSEN 237
BVH11-2 SP64
BVH11-2 JNR7/87 179 DTGDAYIVPHGDHYHYIPKNELSASELAAAEAYWNGKQGSRPSSSSSYNANPAQPRLSEN 238
               179 DTGDAYIVPHGNHFHYIPKSDLSASELAAAQAYWNGKQGSRPSSSSSHNANPAQPRLSEN 238
BVH11-2 P4241
                179 DTGDAYIVPHGNHFHYIPKSDLSASELAAAQAYWNGKQGSRPSSSSSHNANPAQPRLSEN 238
BVH11-2 A66
                179 DTGDAYIVPRGNHFHYIPKSDLSASELAAAQAYWNGKQGSRPSSSSSHNANPAQPRLSEN 238
BVH11-2 WU2
                178 DTGDAYIVPHGDHYHYIPKSDLSASELAAAQAYWNGKQGSRPSSSSSHNANPAQPRLSEN 237
BVH11-2 Rx1
                179 DTGDAYIVPHGNHFHYIPKSDLSASELAAAQAYWNGKQGSRPSSSSSHNANPAQPRLSEN 238
BVH11 P4241
                179 DTGDAYIVPHGNHFHYIPKSDLSASELAAAQAYWNGKQGSRPSSSSSHNANPAQPRLSEN 238
BVH11 WU2
                179 DTGDAYIVPHGNHFHYIPKSDLSASELAAAQAYWNGKQGSRPSSSSSHNANPAQPRLSEN 238
BVH11 A66
                178 DTGDAYIVPHGDHYHYIPKSDLSASELAAAQAYWNGKQGSRPSSSSSHNANPAQPRLSEN 237
BVH11 Rx1
                178 DTGDAYIVPHGDHYHYIPKNELSASELAAAEAYWNGKQGSRPSSSSSYNANPAQPRLSEN 237
BVH11 JNR7/87
                178 DTGDAYIVPHGNHFHYIPKSDLSASELAAAQAYWNGKQGSRPSSSSSHNANPAQPRLSEN 237
BVH11 SP63
                180 DTGDAYIVPHGDHYHYIPKNELSASELAAAEAFLSGRENLSNLRTYRRQNSDNTPRTNWV 239
BVH11 SP64
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238 HNLTVTPTYHQN------QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS 285
BVH11-2 SP64
BVH11-2 JNR7/87 239 HNLTVTPTYHQN------QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS 286
BVH11-2 P4241 239 HNLTVTPTYHQN------QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS 286
               239 HNLTVTPTYHQN------QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS 286
BVH11-2 A66
               239 HNLTVTPTYHQN------QGENISSLLRELYAKPLSERRVESDGLIFDPAQITS 286
BVH11-2 WU2
               238 HNLTVTPTYHQN------QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS 285
BVH11-2 Rx1
               239 HNLTVTPTYHQN------QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS 286
BVH11 P4241
               239 HNLTVTPTYHQN------QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS 286
BVH11 WU2
               239 HNLTVTPTYHQN------QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS 286
BVH11 A66
               238 HNLTVTPTYHQN------QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS 285
BVH11 Rx1
               238 HNLTVTPTYHQN-----QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS 285
BVH11 JNR7/87
               238 HNLTVTPTYHQN------QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS 285
BVH11 SP63
               240 PSVSNPGTTNTNTSNNSNTNSQASQSNDIDSLLKQLYKLPLSQRHVESDGLIFDPAQITS 299
BVH11 SP64
                                             * . *
               286 RTARGVAVPHGNHYHFIPYEQMSELEKRIARIIPLRYRSNHWVPDSRPEQPSPQSTPEPS 345
BVH11-2 SP64
BVH11-2 JNR7/87 287 RTARGVAVPHGNHYHFIPYEQMSELEKRIARIIPLRYRSNHWVPDSRPEQPSPQSTPEPS 346
               287 RTARGVAVPHGNHYHFIPYEQMSELEERIARIIPLRYRSNHWVPDSRPEQPSPQ----PS 342
BVH11-2 P4241
               287 RTARGVAVPHGNHYHFIPYEQMSELEERIARIIPLRYRSNHWVPDSRPEQPSPQ----PS 342
BVH11-2 A66
               287 RTARGVAVPHGNHYHFIPYEQMSELEERIARIIPLRYRSNHWVPDSRPEQPSPQ----PS 342
BVH11-2 WU2
               286 RTANGVAVPHGDHYHFIPYSQLSPLEEKLARIIPLRYRSNHWVPDSRPEQPSPQSTPEPS 345
BVH11-2 Rx1
               287 RTARGVAVPHGNHYHFIPYEQMSELEERIARIIPLRYRSNHWVPDSRPEQPSPQ----PS 342
BVH11 P4241
               287 RTARGVAVPHGNHYHFIPYEQMSELEERIARIIPLRYRSNHWVPDSRPEQPSPQ----PS 342
BVH11 WU2
               287 RTARGVAVPHGNHYHFIPYEQMSELEERIARIIPLRYRSNHWVPDSRPEQPSPQ----PS 342
BVH11 A66
               286 RTANGVAVPHGDHYHFIPYSQLSPLEEKLARIIPLRYRSNHWVPDSRPEQPSPQSTPEPS 345
BVH11 Rx1
               286 RTARGVAVPHGNHYHFIPYEQMSELEKRIARIIPLRYRSNHWVPDSRPEEPSPQPTPEPS 345
BVH11 JNR7/87
               286 RTARGVAVPHGNHYHFIPYSQMSELEERIARIIPLRYRSNHWVPDSRPEQPSPQSTPEPS 345
BVH11 SP63
               300 RTARGVAVPHGNHYHFIPYEQMSELEKRIARIIPLRYRSNHWVPDSRPEEPSPQPTPEPS 359
BVH11 SP64
                   346 PSLQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYIPAKDLSAETAAGIDSK 405
BVH11-2 SP64
BVH11-2 JNR7/87 347 PSPQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYIPAKDLSAETAAGIDSK 406
               343 PSPQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYIPAKDLSAETAAGIDSK 402
BVH11-2 P4241
               343 PSPQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYIPAKDLSAETAAGIDSK 402
BVH11-2 A66
               343 PSPQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYIPAKDLSAETAAGIDSK 402
BVH11-2 WU2
               346 PSPQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVPRYIPAKDLSAETAAGIDSK 405
BVH11-2 Rx1
               343 PSPQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYIPAKDLSAETAAGIDSK 402
BVH11 P4241
               343 PSPQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYIPAKDLSAETAAGIDSK 402
BVH11 WU2
               343 PSPQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYIPAKDLSAETAAGIDSK 402
BVH11 A66
               346 PSPQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVPRYIPAKDLSAETAAGIDSK 405
BVH11 Rx1
               346 PSP-----QPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYIPAKDLSAETAAGIDSK 399
BVH11 JNR7/87
               346 PSPQSAPNPQPAPSNPIDEKLVKEVVRKVGDGYVFEKNGVSRYIPAKNLSAETAAGIDSK 405
BVH11 SP63
               360 PSPQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYIPAKNLSAETAAGIDSK 419
BVH11 SP64
                            *************
               406 LAKQESLSHKLGAKKTDLPSSDREFYNKAYDLLARIHQDLLDNKGRQVDFEVLDNLLERL 465
BVH11-2 SP64
BVH11-2 JNR7/87 407 LAKQESLSHKLGAKKTDLPSSDREFYNKAYDLLARIHQDLLDNKGRQVDFEALDNLLERL 466
               403 LAKQESLSHKLGTKKTDLPSSDREFYNKAYDLLARIHQDLLDNKGRQVDFEALDNLLERL 462
BVH11-2 P4241
               403 LAKQESLSHKLGTKKTDLPSSDREFYNKAYDLLARIHQDLLDNKGRQVDFEALDNLLERL 462
BVH11-2 A66
               403 LAKQESLSHKLGTKKTDLPSSDREFYNKAYDLLARIHQDLLDNKGRQVDFEALDNLLERL 462
BVH11-2 WU2
               406 LAKQESLSHKLGAKKTDLPSSDREFYNKAYDLLARIHQDLLDNKGRQVDFEALDNLLERL 465
BVH11-2 Rx1
               403 LAKQESLSHKLGTKKTDLPSSDREFYNKAYDLLARIHQDLLDNKGRQVDFEALDNLLERL 462
BVH11 P4241
               403 LAKQESLSHKLGTKKTDLPSSDREFYNKAYDLLARIHQDLLDNKGRQVDFEALDNLLERL 462
BVH11 WU2
               403 LAKQESLSHKLGTKKTDLPSSDREFYNKAYDLLARIHQDLLDNKGRQVDFEALDNLLERL 462
BVH11 A66
               406 LAKQESLSHKLGAKKTDLPSSDREFYNKAYDLLARIHQDLLDNKGRQVDFEALDNLLERL 465
BVH11 Rx1
               400 LAKQESLSHKLGAKKTDLPSSDREFYNKAYDLLARIHQDLLDNKGRQVDFEALDNLLERL 459
BVH11 JNR7/87
               406 LAKQESLSHKLGAKKTDLPSSDREFYNKAYDLLARIHQDLLDNKGRQVDFEALDNLLERL 465
BVH11 SP63
               420 LAKQESLSHKLGAKKTDLPSSDREFYNKAYDLLARIHQDLLDNKGRQVDFEALDNLLERL 479
BVH11 SP64
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BVH11-2 SP64
                466 KDVSSDKVKLVDDILAFLAPIRHPERLGKPNAQITYTDDEIQVAKLAGKYTTEDGYIFDP 525
BVH11-2 JNR7/87 467 KDVPSDKVKLVDDILAFLAPIRHPERLGKPNAQITYTDDEIQVAKLAGKYTTEDGYIFDP 526
BVH11-2 P4241 463 KDVSSDKVKLVEDILAFLAPIRHPERLGKPNSQITYTDDEIQVAKLAGKYTTEDGYIFDP 522
                463 KDVSSDKVKLVEDILAFLAPIRHPERLGKPNSQITYTDDEIQVAKLAGKYTTEDGYIFDP 522
BVH11-2 A66
BVH11-2 WU2
               463 KDVSSDKVKLVEDILAFLAPIRHPERLGKPNSQITYTDDEIQVAKLAGKYTTEDGYIFDP 522
               466 KDVSSDKVKLVDDILAFLAPIRHPERLGKPNAQITYTDDEIQVAKLAGKYTTEDGYIFDP 525
BVH11-2 Rx1
BVH11 P4241
               463 KDVSSDKVKLVEDILAFLAPIRHPERLGKPNSQITYTDDEIQVAKLAGKYTTEDGYIFDP 522
BVH11 WU2
               463 KDVSSDKVKLVEDILAFLAPIRHPERLGKPNSQITYTDDEIQVAKLAGKYTTEDGYIFDP 522
BVH11 A66
               463 KDVSSDKVKLVEDILAFLAPIRHPERLGKPNSQITYTDDEIQVAKLAGKYTTEDGYIFDP 522
               466 KDVSSDKVKLVDDILAFLAPIRHPERLGKPNAQITYTDDEIQVAKLAGKYTTEDGYIFDP 525
BVH11 Rx1
BVH11 JNR7/87
               460 KDVSSDKVKLVDDILAFLAPIRHPERLGKPNAQITYTDDEIQVAKLAGKYTTEDGYIFDP 519
BVH11 SP63
               466 EDVPSDKVKLVDDILAFLAPIRHPERLGKPNAQITYTDDEIQVAKLAGKYTTEDGYIFDP 525
               480 KDVSSDKVKLVDDILAFLAPIRHPERLGKPNAQITYTDDEIQVAKLAGKYTTEDGYIFDP 539
                    ** ****** . ************* . **********
BVH11-2 SP64
               526 RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHODSGNTEAK 585
BVH11-2 JNR7/87 527 RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK 586
BVH11-2 P4241
               523 RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHRDSGNTEAK 582
BVH11-2 A66
               523 RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHODSGNTEAK 582
BVH11-2 WU2
               523 RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK 582
BVH11-2 Rx1
               526 RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAOAYAKEKGLTPPSTDHODSGNTEAK 585
BVH11 P4241
               523 RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK 582
BVH11 WU2
               523 RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK 582
               523 RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK 582
BVH11 A66
BVH11 Rx1
               526 RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHODSGNTEAK 585
BVH11 JNR7/87
               520 RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK 579
BVH11 SP63
               526 RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHODSGNTEAK 585
BVH11 SP64
               540 RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK 599
                   ******************
BVH11-2 SP64
               586 GAEAIYNRVKAAKKVPLDRMPYNLQYTVEVKNGSLIIPHYDHYHNIKFEWFDEGLYEAPK 645
BVH11-2 JNR7/87 587 GAEAIYNRVKAAKKVPLDRMPYNLQYTVEVKNGSLIIPHYDHYHNIKFEWFDEGLYEAPK 646
BVH11-2 P4241
               583 GAEAIYNRVKAAKKVPLDRMPYNLQYTVEVKNGSLIIPHYDHYHNIKFEWFDEGLYEAPK 642
               583 GAEAIYNRVKAAKKVPLDRMPYNLQYTVEVKNGSLIIPHYDHYHNIKFEWFDEGLYEAPK 642
BVH11-2 A66
BVH11-2 WU2
               583 GAEAIYNRVKAAKKVPLDRMPYNLQYTVEVKNGSLIIPHYDHYHNIKFEWFDEGLYEAPK 642
BVH11-2 Rx1
               586 GAEAIYNRVKAAKKVPLDRMPYNLQYTVEVKNGSLIIPHYDHYHNIKFEWFDEGLYEAPK 645
BVH11 P4241
               583 GAEAIYNRVKAAKKVPLDRMPYNLQYTVEVKNGSLIIPHYDHYHNIKFEWFDEGLYEAPK 642
BVH11 WU2
               583 GAEAIYNRVKAAKKVPLDRMPYNLQYTVEVKNGSLIIPHYDHYHNIKFEWFDEGLYEAPK 642
BVH11 A66
               583 GAEAIYNRVKAAKKVPLDRMPYNLQYTVEVKNGSLIIPHYDHYHNIKFEWFDEGLYEAPK 642
BVH11 Rx1
               586 GAEAIYNRVKAAKKVPLDRMPYNLQYTVEVKNGSLIIPHYDHYHNIKFEWFDEGLYEAPK 645
BVH11 JNR7/87
               580 GAEAIYNRVKAAKKVPLDRMPYNLQYTVEVKNGSLIIPHYDHYHNIKFEWFDEGLYEAPK 639
BVH11 SP63
               586 GAEAIYNRVKAAKKVPLDRMPYNLQYTVEVKNGSLIIPHYDHYHNIKFEWFDEGLYEAPK 645
BVH11 SP64
               600 GAEAIYNRVKAAKKVPLDRMPYNLQYTVEVKNGSLIIPHYDHYHNIKFEWFDEGLYEAPK 659
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               646 GYSLEDLLATVKYYVEHPNERPHSDNGFGNASDHVRKNK------ADQDSK 690
BVH11-2 SP64
BVH11-2 JNR7/87 647 GYTLEDLLATVKYYVEHPNERPHSDNGFGNASDHVRKNK------VDQDSK 691
BVH11-2 P4241
               643 GYTLEDLLATVKYYVEHPNERPHSDNGFGNASDHVRKNK------ADODSK 687
BVH11-2 A66
               643 GYTLEDLLATVKYYVEHPNERPHSDNGFGNASDHVRKNK------ADODSK 687
BVH11-2 WU2
               643 GYTLEDLLATVKYYVEHPNERPHSDNGFGNASDHVRKNK------ADODSK 687
BVH11-2 Rx1
               646 GYSLEDLLATVKYYVEHPNERPHSDNGFGNASDHVQRNKNGQADTNQTEKPNEEKPQTEK 705
BVH11 P4241
               643 GYTLEDLLATVKYYVEHPNERPHSDNGFGNASDHVRKNK------ADODSK 687
               643 GYTLEDLLATVKYYVEHPNERPHSDNGFGNASDHVRKNK------ADQDSK 687
BVH11 WU2
BVH11 A66
               643 GYTLEDLLATVKYYVEHPNERPHSDNGFGNASDHVRKNK------ADODSK 687
               646 GYSLEDLLATVKYYVEHPNERPHSDNGFGNASDHVQRNK-----NGQ 687
BVH11 Rx1
BVH11 JNR7/87
              640 GYSLEDLLATVKYYVEHPNERPHSDNGFGNASDHVQRNK-----NGQ 681
BVH11 SP63
               646 GYTLEDLLATVKYYVEHPNERPHSDNGFGNASDHVQRNK-----NGO 687
               660 GYTLEDLLATVKYYVEHPNERPHSDNGFGNASDHVQRNK-----NGQ 701
BVH11 SP64
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BVH11-2 SP64
                691 PDEDKEHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEAEDTTDEAEIPOV 750
BVH11-2 JNR7/87 692 PDEDKEHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEAEDTTDEAEIPQV 751
BVH11-2 P4241
                688 PDEDKGHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEAEDTTDEAEIPQV 747
                688 PDEDKGHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEAEDTTDEAEIPQV 747
BVH11-2 A66
BVH11-2 WU2
                688 PDEDKGHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEAEDTTDEAEIPQV 747
                706 PEEDKEHDEVSEPTHPESDEKENHVGLNPSADNLYKPSTDTEETEEEAEDTTDEAEIPQV 765
BVH11-2 Rx1
BVH11 P4241
                688 PDEDKGHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEAEDTTDEAEIPQV 747
BVH11 WU2
                688 PDEDKGHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEAEDTTDEAEIPQV 747
BVH11 A66
                688 PDEDKGHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEAEDTTDEAEIPQV 747
                688 ADTNQTEKPNEEKPQTEKPEEETPREEKPQSEKPESPKPTEEPEEESPEESEEPQV 747
BVH11 Rx1
BVH11 JNR7/87
                682 ADTNQTEKPNEEKPQTEKPEEETPREEKPQSEKPESPKPTEEPEEESPEESPEESEEPQV 741
                688 ADTNQTEKPSEEKPQTEKPEEETPREEKPQSEKPESP----KPTEEPEEESPEESEEPQV 743
BVH11 SP63
BVH11 SP64
                702 ADTNQTEKPSEEKPQTEKPEEETPREEKPQSEKPESP----KPTEEPEEESPEESEEPQV 757
                                                * ..
                               * . * * *.
BVH11-2 SP64
                751 ENSVINAKIADAEALLEKVTDPSIRQNAMETLTGLKSSLLLGTKDNNTISAEVDSLLALL 810
BVH11-2 JNR7/87 752 ENSVINAKIADAEALLEKVTDPSIRQNAMETLTGLKSSLLLGTKDNNTISAEVDSLLALL 811
BVH11-2 P4241
                748 EHSVINAKIADAEALLEKVTDPSIRQNAMETLTGLKSSLLLGTKDNNTISAEVDSLLALL 807
BVH11-2 A66
                748 EHSVINAKIADAEALLEKVTDPSIRQNAMETLTGLKSSLLLGTKDNNTISAEVDSLLALL 807
BVH11-2 WU2
                748 EHSVINAKIADAEALLEKVTDPSIRQNAMETLTGLKSSLLLGTKDNNTISAEVDSLLALL 807
BVH11-2 Rx1
                766 EYSVINAKIAEAEALLEKVTDSSIRQNAVETLTGLKSSLLLGTKDNNTISAEVDSLLALL 825
BVH11 P4241
                748 EHSVINAKIADAEALLEKVTDPSIRQNAMETLTGLKSSLLLGTKDNNTISAEVDSLLALL 807
BVH11 WU2
                748 EHSVINAKIADAEALLEKVTDPSIRQNAMETLTGLKSSLLLGTKDNNTISAEVDSLLALL 807
BVH11 A66
                748 EHSVINAKIADAEALLEKVTDPSIRQNAMETLTGLKSSLLLGTKDNNTISAEVDSLLALL 807
BVH11 Rx1
                748 ETEKVKEKLREAEDLLGKIQNPIIKSNAKETLTGLKNNLLFGTQDNNTIMAEAEKLLALL 807
BVH11 JNR7/87
                742 ETEKVKEKLREAEDLLGKIQNPIIKSNAKETLTGLKNNLLFGTQDNNTIMAEAEKLLALL 801
BVH11 SP63
                744 ETEKVEEKLREAEDLLGKIQDPIIKSNAKETLTGLKNNLLFGTQDNNTIMAEAEKLLALL 803
BVH11 SP64
                758 ETEKVEEKLREAEDLLGKIQDPIIKSNAKETLTGLKNNLLFGTQDNNTIMAEAEKLLALL 817
                        . *. .** ** *.
                                          *. ** ***** ** ** **** ** . ****
BVH11-2 SP64
                811 KESQPAPIQ 819
BVH11-2 JNR7/87 812 KESQPAPIQ 820
BVH11-2 P4241
               808 KKSQPAPIQ 816
BVH11-2 A66
                808 KKSQPAPIQ 816
BVH11-2 WU2
                808 KKSOPAPIO 816
BVH11-2 Rx1
              826 KESQPAPIQ 834
BVH11 P4241
                808 KESK
                              811
BVH11 WU2
                808 KESK
                              811
BVH11 A66
                808 KESK
                              811
BVH11 Rx1
                808 KESK
                              811
BVH11 JNR7/87
                802 KESK
                              805
BVH11 SP63
                804 KESK
                              807
BVH11 SP64
                818 KESK
                              821
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FIGURE 12

	[7				Ī	7	3	7			7				7		Γ	-	7			
	BVH11	SP64	BVH11-2	SP64	BVH11	SP63	BVH11	JNR.7/87	BVH11-2	JNR.7/87	BVH11	WU2	BVH11-2	WU2	BVH11	A66	BVH11-2	99Y	BVH11	P4241	BVH11-2	P4241	BVH11	Rx-1
BVH11-2 Rx-1	I 81%	S 85%	I 94%	S 95%	%68 I	S 91%	%88 I	%06 S	I 94%	S 95%	1 92%	S 94%	I 93%	S 95%	I 92%	S 94%	I 93%	S 95%	I 92%	S 94%	I 93%	S 95%	191%	S 92%
BVH11 Rx-1	%88 I	S 91%	I 87%	%06 S	%26 I	%26 S	%96 I	%96 S	%28 I	%06 S	I 87%	S 91%	%98 I	%06 S	%L8 I	S 91%	%98 I	%06 S	% <i>L</i> 8 I	S 91%	%98 I	%06 S		
BVH11-2 BVH1 P4241 Rx-1	%08 I	82% S	%96 I	% 26 S	%/8 I	%06 S	%98 I	%06 S	%26 I	%86 S	%86 I	%86 S	%66 I	%66 S	%66 I	%66 S	%66 I	%66 S	%66 I	%66 S				
BVH11 P4241	%08 I	S 85%	%56 I	%96 S	%88 I	S 91%	%L8 I	S 91%	%96 I	8 97%	%66 I	%66 S	%86 I	%86 S	I 100%		%66 I	%66 S						
BVH11-2 A66	%08 I	S 85%	%96 I	%16 S	%L8 I	%06 S	%98 I	%06 S	% <i>L</i> 6 I	%86 S	%86 I	%86 S	%66 I	%66 S	%66 I	%66 S								
BVH11 A66	%08 I	S 85%	%56 I	%96 S	%88 I	S 91%	%L8 I	S 91%	%96 I	%26 S	I 92%	S 94%	%86 I	%86 S										
BVH11-2 WU2	%08 I	S 85%	%96 I	8 97%	I 87%	%06 S	%98 I	%06 S	%/6 I	%86 S	%86 I	%86 S												
BVH11 WU2	%08 I	S 85%	%56 I	%96 S	%88 I	S 91%	%L8 I	S 91%	%96 I	S 97%														
	I 82%	S 87%	%86 I	%86 S	%88 I	S 91%	%L8 I	%06 S													13			
BVH11 JNR.7/87	%88 I	S 91%	%/8 I	%06 S	%96 I	%96 S															FIGURE 13			
BVH11 SP63	%88 I	%06 S	%L8 I	%06 S																				
BVH11-2 BVH11 SP64 SP63	I 81%	%98 S																						

AATTCCTTGT CGGGTAAGTT CCGACCCGCA CGAAAGGCGT AATGATTTGG GCACTGTCTC 60 AACGAGAGAC TCGGTGAAAT TTTAGTACCT GTGAAGATGC AGGTTACCCG CGACAGGACG 120 GAAAGACCCC ATGGAGCTTT ACTGCAGTTT GATATTGAGT GTCTGTACCA CATGTACAGG 180 ATAGGTAGGA GTCTAAGAGA TCGGGACGCC AGTTTCGAAG GAGACGCTGT TGGGATACTA 240 CCCTTGTGTT ATGGCCACTC TAACCCAGAT AGGTGATCCC TATCGGAGAC AGTGTCTGAC 300 GGGCAGTTTG ACTGGGGCGG TCGCCTCCTA AAAGGTAACG GAGGCGCCCA AAGGTTCCCT 360 CAGAATGGTT GGAAATCATT CGCAGAGTGT AAAGGTATAA GGGAGCTTGA CTGCGAGAGC 420 TACAACTCGA GCAGGGACGA AAGTCGGGCT TAGTGATCCG GTGGTTCCGT ATGGAAGGGC 480 CATCGCTCAA CGGATAAAAG CTACCCTGGG GATAACAGGC TTATCTCCCC CAAGAGTTCA 540 CATCGACGGG GAGGTTTGGC ACCTCGATGT CGGCTCGTCG CATCCTGGGG CTGTAGTCGG 600 TCCCAAGGGT TGGGCTGTTC GCCCATTAAA GCGGCACGCG AGCTGGGTTC AGAACGTCGT 660 GAGACAGTTC GGTCCCTATC CGTCGCGGGC GTAGGAAATT TGAGAGGATC TGCTCCTAGT 720 ACGAGAGGAC CAGAGTGGAC TTACCGCTGG TGTACCAGTT GTCTTGCCAA AGGCATCGCT 780 GGGTAGCTAT GTAGGGAAGG GATAAACGCT GAAAGCATCT AAGTGTGAAA CCCACCTCAA 840 GATGAGATTT CCCATGATTA TATATCAGTA AGAGCCCTGA GAGATGATCA GGTAGATAGG 900 TTAGAAGTGG AAGTGTGGCG ACACATGTAG CGGACTAATA CTAATAGCTC GAGGACTTAT 960 CCAAAGTAAC TGAGAATATG AAAGCGAACG GTTTTCTTAA ATTGAATAGA TATTCAATTT 1020 TGAGTAGGTA TTACTCAGAG TTAAGTGACG ATAGCCTAGG AGATACACCT GTACCCATGC 1080 CGAACACAGA AGTTAAGCCC TAGAACGCCG GAAGTAGTTG GGGGTTGCCC CCTGTGAGAT 1140 AGGGAAGTCG CTTAGCTCTA GGGAGTTTAG CTCAGCTGGG AGAGCATCTG CCTTACAAGC AGAGGGTCAG CGGTTCGATC CCGTTAACTC CCAAAGGTCC CGTAGTGTAG CGGTTATCAC 1260 GTCGCCCTGT CACGGCGAAG ATCGCGGGTT CGATTCCCGT CGGGACCGTT TAAGGTAACG 1320 CAAGTTATTT TAGACTCGTT AGCTCAGTTG GTAGAGCAAT TGACTTTTAA TCAATGGGTC 1380 ACTGGTTCGA GCCCAGTACG GGTCATATAT GCGGGTTTGG CGGAATTCTA ATCTCTTTGA 1440 AATCATCTTC TCTCACTTTC CAAAACTCTA TTACCTCTTA TTATACCACA TTTCAATCTT 1500 CAACTTCCCA GTAATATAAG CACCTCTGGC GAAAGAAGTT TCAATGTCCT AAAGTAATAA 1560 GTGAATCCAA TTCAGGAACT CCAAGAACAA AAGAAACATC TGGTGTCACA AGTATTGGAT GGCACAGAGT CACGTGGTAG TCTGACCCTA GCAGAAATTT TAAATAGTAA ACTATTTACT 1680 GGTTAATTAA ATGGTTAAAT AACCGGTTTA GAAAACTATT TAATAAAGTA AAAGAAGTTG 1740 AGAAAAACT TCATCATTTA TTGAAATGAG GGATTTATGA AATTTAGTAA AAAATATATA 1800 GCAGCTGGAT CAGCTGTTAT CGTATCCTTG AGTCTATGTG CCTATGCACT AAACCAGCAT 1860 CGTTCGCAGG AAAATAAGGA CAATAATCGT GTCTCTTATG TGGATGGCAG CCAGTCAAGT 1920 CAGAAAAGTG AAAACTTGAC ACCAGACCAG GTTAGCCAGA AAGAAGGAAT TCAGGCTGAG 1980 CAAATTGTAA TCAAAATTAC AGATCAGGGC TATGTAACGT CACACGGTGA CCACTATCAT TACTATAATG GGAAAGTTCC TTATGATGCC CTCTTTAGTG AAGAACTCTT GATGAAGGAT CCAAACTATC AACTTAAAGA CGCTGATATT GTCAATGAAG TCAAGGGTGG TTATATCATC 2160 AAGGTCGATG GAAAATATTA TGTCTACCTG AAAGATGCAG CTCATGCTGA TAATGTTCGA 2220 ACTAAAGATG AAATCAATCG TCAAAAACAA GAACATGTCA AAGATAATGA GAAGGTTAAC TCTAATGTTG CTGTAGCAAG GTCTCAGGGA CGATATACGA CAAATGATGG TTATGTCTTT 2340 AATCCAGCTG ATATTATCGA AGATACGGGT AATGCTTATA TCGTTCCTCA TGGAGGTCAC 2400 TATCACTACA TTCCCAAAAG CGATTTATCT GCTAGTGAAT TAGCAGCAGC TAAAGCACAT CTGGCTGGAA AAAATATGCA ACCGAGTCAG TTAAGCTATT CTTCAACAGC TAGTGACAAT 2520 AACACGCAAT CTGTAGCAAA AGGATCAACT AGCAAGCCAG CAAATAAATC TGAAAATCTC 2580 CAGAGTCTTT TGAAGGAACT CTATGATTCA CCTAGCGCCC AACGTTACAG TGAATCAGAT 2640 GGCCTGGTCT TTGACCCTGC TAAGATTATC AGTCGTACAC CAAATGGAGT TGCGATTCCG 2700 CATGGCGACC ATTACCACTT TATTCCTTAC AGCAAGCTTT CTGCTTTAGA AGAAAAGATT 2760 GCCAGAATGG TGCCTATCAG TGGAACTGGT TCTACAGTTT CTACAAATGC AAAACCTAAT 2820 GAAGTAGTGT CTAGTCTAGG CAGTCTTTCA AGCAATCCTT CTTCTTTAAC GACAAGTAAG GAGCTCTCTT CAGCATCTGA TGGTTATATT TTTAATCCAA AAGATATCGT TGAAGAAACG 3000 ATTGGGCAAC CGACTCTTCC AAACAATAGT CTAGCAACAC CTTCTCCATC TCTTCCAATC 3060 AATCCAGGAA CTTCACATGA GAAACATGAA GAAGATGGAT ACGGATTTGA TGCTAATCGT 3120 ATTATCGCTG AAGATGAATC AGGTTTTGTC ATGAGTCACG GAGACCACAA TCATTATTTC 3180 TTCAAGAAGG ACTTGACAGA AGAGCAAATT AAGGCTGCGC AAAAACATTT AGAGGAAGTT 3240 AAAACTAGTC ATAATGGATT AGATTCTTTG TCATCTCATG AACAGGATTA TCCAGGTAAT 3300 GCCAAAGAAA TGAAAGATTT AGATAAAAAA ATCGAAGAAA AAATTGCTGG CATTATGAAA

CAATATGGTG TCAAACGTGA AAGTATTGTC GTGAATAAAG AAAAAAATGC GATTATTTAT 3420 CCGCATGGAG ATCACCATCA TGCAGATCCG ATTGATGAAC ATAAACCGGT TGGAATTGGT CATTCTCACA GTAACTATGA ACTGTTTAAA CCCGAAGAAG GAGTTGCTAA AAAAGAAGGG 3540 AATAAAGTTT ATACTGGAGA AGAATTAACG AATGTTGTTA ATTTGTTAAA AAATAGTACG 3600 TTTAATAATC AAAACTTTAC TCTAGCCAAT GGTCAAAAAC GCGTTTCTTT TAGTTTTCCG 3660 CCTGAATTGG AGAAAAATT AGGTATCAAT ATGCTAGTAA AATTAATAAC ACCAGATGGA 3720 AAAGTATTGG AGAAAGTATC TGGTAAAGTA TTTGGAGAAG GAGTAGGGAA TATTGCAAAC 3780 TTTGAATTAG ATCAACCTTA TTTACCAGGA CAAACATTTA AGTATACTAT CGCTTCAAAA GATTATCCAG AAGTAAGTTA TGATGGTACA TTTACAGTTC CAACCTCTTT AGCTTACAAA 3900 ATGGCCAGTC AAACGATTTT CTATCCTTTC CATGCAGGGG ATACTTATTT AAGAGTGAAC 3960 CCTCAATTTG CAGTGCCTAA AGGAACTGAT GCTTTAGTCA GAGTGTTTGA TGAATTTCAT 4020 GGAAATGCTT ATTTAGAAAA TAACTATAAA GTTGGTGAAA TCAAATTACC GATTCCGAAA 4080 TTAAACCAAG GAACAACCAG AACGGCCGGA AATAAAATTC CTGTAACCTT CATGGCAAAT 4140 GCTTATTTGG ACAATCAATC GACTTATATT GTGGAAGTAC CTATCTTGGA AAAAGAAAAT 4200 CAAACTGATA AACCAAGTAT TCTACCACAA TTTAAAAGGA ATAAAGCACA AGAAAACTCA 4260 AAACTTGATG AAAAGGTAGA AGAACCAAAG ACTAGTGAGA AGGTAGAAAA AGAAAAACTT 4320 TCTGAAACTG GGAATAGTAC TAGTAATTCA ACGTTAGAAG AAGTTCCTAC AGTGGATCCT 4380 GTACAAGAAA AAGTAGCAAA ATTTGCTGAA AGTTATGGGA TGAAGCTAGA AAATGTCTTG 4440 TTTAATATGG ACGGAACAAT TGAATTATAT TTACCATCAG GAGAAGTCAT TAAAAAGAAT 4500 ATGGCAGATT TTACAGGAGA AGCACCTCAA GGAAATGGTG AAAATAAACC ATCTGAAAAT 4560 GGAAAAGTAT CTACTGGAAC AGTTGAGAAC CAACCAACAG AAAATAAACC AGCAGATTCT TTACCAGAGG CACCAAACGA AAAACCTGTA AAACCAGAAA ACTCAACGGA TAATGGAATG 4680 TTGAATCCAG AAGGGAATGT GGGGAGTGAC CCTATGTTAG ATCCAGCATT AGAGGAAGCT 4740 CCAGCAGTAG ATCCTGTACA AGAAAAATTA GAAAAATTTA CAGCTAGTTA CGGATTAGGC 4800 TTAGATAGTG TTATATTCAA TATGGATGGA ACGATTGAAT TAAGATTGCC AAGTGGAGAA 4860 GTGATAAAAA AGAATTTATC TGATTTCATA GCGTAAGGAA TAGCAGTAGA AAAAGTCTGA 4920 ATCAAAAATG AAGTTCTCTC AAAAGTTAGA AATAAAACTC TGACTTTGGG AGAATTTCAT 4980 TTTATTATTA ATATAAAA TTTCTTGACA TACAACTTAA AAAGAGGTGG AATATTTACT 5040 AGTTAATT (SEQ ID NO : 11) 5048

FIGURE 14

CAGAGATCTT AGTGAATCAA ATATACTTAA GAAAAGAGGA AAGAATGAAA ATCAATAAAA 60 AATATCTAGC TGGGTCAGTA GCTACACTTG TTTTAAGTGT CTGTGCTTAT GAACTAGGTT 120 TGCATCAAGC TCAAACTGTA AAAGAAAATA ATCGTGTTTC CTATATAGAT GGAAAACAAG 180 CGACGCAAAA AACGGAGAAT TTGACTCCTG ATGAGGTTAG CAAGCGTGAA GGAATCAACG 240 CCGAACAAAT CGTCATCAAG ATTACGGATC AAGGTTATGT GACCTCTCAT GGAGACCATT 300 ATCATTACTA TAATGGCAAG GTCCCTTATG ATGCCATCAT CAGTGAAGAG CTCCTCATGA 360 AAGATCCGAA TTATCAGTTG AAGGATTCAG ACATTGTCAA TGAAATCAAG GGTGGTTATG 420 TCATTAAGGT AAACGGTAAA TACTATGTTT ACCTTAAGGA TGCAGCTCAT GCGGATAATG TCCGTACAAA AGAAGAAATC AATCGGCAAA AACAAGAACA TAGTCAGCAT CGTGAAGGAG 540 GGACTTCAGC AAACGATGGT GCGGTAGCCT TTGCACGTTC ACAGGGACGC TACACCACAG 600 ATGATGGTTA TATCTTCAAT GCATCTGATA TCATCGAAGA TACGGGCGAT GCCTATATCG 660 TTCCTCATGG AGATCATTAC CATTACATTC CTAAGAATGA GTTATCAGCT AGCGAGTTGG 720 CTGCTGCAGA AGCCTTCCTA TCTGGTCGGG AAAATCTGTC AAATTTAAGA ACCTATCGCC 780 GACAAAATAG CGATAACACT CCAAGAACAA ACTGGGTACC TTCTGTAAGC AATCCAGGAA 840 CTACAAATAC TAACACAAGC AACAACAGCA ACACTAACAG TCAAGCAAGT CAAAGTAATG 900 ACATTGATAG TCTCTTGAAA CAGCTCTACA AACTGCCTTT GAGTCAACGC CATGTAGAAT 960 CTGATGGCCT TATTTTCGAC CCAGCGCAAA TCACAAGTCG AACCGCCAGA GGTGTAGCTG 1020 TCCCTCATGG TAACCATTAC CACTTTATCC CTTATGAACA AATGTCTGAA TTGGAAAAAC 1080 GAATTGCTCG TATTATTCCC CTTCGTTATC GTTCAAACCA TTGGGTACCA GATTCAAGAC 1140 CAGAAGAACC AAGTCCACAA CCGACTCCAG AACCTAGTCC AAGTCCGCAA CCTGCACCAA 1200 ATCCTCAACC AGCTCCAAGC AATCCAATTG ATGAGAAATT GGTCAAAGAA GCTGTTCGAA AAGTAGGCGA TGGTTATGTC TTTGAGGAGA ATGGAGTTTC TCGTTATATC CCAGCCAAGA 1320 ATCTTTCAGC AGAAACAGCA GCAGGCATTG ATAGCAAACT GGCCAAGCAG GAAAGTTTAT 1380 CTCATAAGCT AGGAGCTAAG AAAACTGACC TCCCATCTAG TGATCGAGAA TTTTACAATA 1440 AGGCTTATGA CTTACTAGCA AGAATTCACC AAGATTTACT TGATAATAAA GGTCGACAAG 1500 TTGATTTTGA GGCTTTGGAT AACCTGTTGG AACGACTCAA GGATGTCTCA AGTGATAAAG 1560 TCAAGTTAGT GGATGATATT CTTGCCTTCT TAGCTCCGAT TCGTCATCCA GAACGTTTAG 1620 GAAAACCAAA TGCGCAAATT ACCTACACTG ATGATGAGAT TCAAGTAGCC AAGTTGGCAG 1680 GCAAGTACAC AACAGAAGAC GGTTATATCT TTGATCCTCG TGATATAACC AGTGATGAGG 1740 GGGATGCCTA TGTAACTCCA CATATGACCC ATAGCCACTG GATTAAAAAA GATAGTTTGT 1800 CTGAAGCTGA GAGAGCGGCA GCCCAGGCTT ATGCTAAAGA GAAAGGTTTG ACCCCTCCTT 1860 CGACAGACCA TCAGGATTCA GGAAATACTG AGGCAAAAGG AGCAGAAGCT ATCTACAACC 1920 GCGTGAAAGC AGCTAAGAAG GTGCCACTTG ATCGTATGCC TTACAATCTT CAATATACTG 1980 TAGAAGTCAA AAACGGTAGT TTAATCATAC CTCATTATGA CCATTACCAT AACATCAAAT TTGAGTGGTT TGACGAAGGC CTTTATGAGG CACCTAAGGG GTATACTCTT GAGGATCTTT TGGCGACTGT CAAGTACTAT GTCGAACATC CAAACGAACG TCCGCATTCA GATAATGGTT TTGGTAACGC TAGCGACCAT GTTCAAAGAA ACAAAAATGG TCAAGCTGAT ACCAATCAAA 2220 CGGAAAAACC AAGCGAGGAG AAACCTCAGA CAGAAAAACC TGAGGAAGAA ACCCCTCGAG AAGAGAAACC ACAAAGCGAG AAACCAGAGT CTCCAAAACC AACAGAGGAA CCAGAAGAAG 2340 AATCACCAGA GGAATCAGAA GAACCTCAGG TCGAGACTGA AAAGGTTGAA GAAAAACTGA 2400 GAGAGGCTGA AGATTTACTT GGAAAAATCC AGGATCCAAT TATCAAGTCC AATGCCAAAG 2460 AGACTCTCAC AGGATTAAAA AATAATTTAC TATTTGGCAC CCAGGACAAC AATACTATTA 2520 TGGCAGAAGC TGAAAAACTA TTGGCTTTAT TAAAGGAGAG TAAGTAAAGG TAGCAGCATT 2580 TTCTAACTCC TAAAAACAGG ATAGGAGAAC GGGAAAACGA AAAATGAGAG CAGAATGTGA 2640 GTTCTAG (SED ID NO : 12) 2647

GGGTCTTAAA ACTCTGAATC CTTTAGAGGC AGACCCACAA AATGACAAGA CCTATTTAGA 60 AAATCTGGAA GAAAATATGA GTGTTCTAGC AGAAGAATTA AAGTGAGGAA AGAATGAAAA 120 TCAATAAAAA ATATCTAGCA GGTTCAGTGG CAGTCCTTGC CCTAAGTGTT TGTTCCTATG 180 AACTTGGTCG TCACCAAGCT GGTCAGGTTA AGAAAGAGTC TAATCGAGTT TCTTATATAG 240 ATGGTGATCA GGCTGGTCAA AAGGCAGAAA ATTTGACACC AGATGAAGTC AGTAAGAGAG 300 AGGGGATCAA CGCCGAACAA ATTGTTATCA AGATTACGGA TCAAGGTTAT GTGACCTCTC 360 ATGGAGACCA TTATCATTAC TATAATGGCA AGGTTCCTTA TGATGCCATC ATCAGTGAAG 420 AACTTCTCAT GAAAGATCCG AATTATCAGT TGAAGGATTC AGACATTGTC AATGAAATCA 480 AGGGTGGCTA TGTGATTAAG GTAGACGGAA AATACTATGT TTACCTTAAA GATGCGGCCC 540 ATGCGGACAA TATTCGGACA AAAGAAGAGA TTAAACGTCA GAAGCAGGAA CACAGTCATA 600 ATCATAACTC AAGAGCAGAT AATGCTGTTG CTGCAGCCAG AGCCCAAGGA CGTTATACAA 660 CGGATGATGG GTATATCTTC AATGCATCTG ATATCATTGA GGACACGGGT GATGCTTATA 720 TCGTTCCTCA CGGCGACCAT TACCATTACA TTCCTAAGAA TGAGTTATCA GCTAGCGAGT 780 TAGCTGCTGC AGAAGCCTAT TGGAATGGGA AGCAGGGATC TCGTCCTTCT TCAAGTTCTA 840 GTTATAATGC AAATCCAGTT CAACCAAGAT TGTCAGAGAA CCACAATCTG ACTGTCACTC 900 CAACTTATCA TCAAAATCAA GGGGAAAACA TTTCAAGCCT TTTACGTGAA TTGTATGCTA 960 AACCCTTATC AGAACGCCAT GTAGAATCTG ATGGCCTTAT TTTCGACCCA GCGCAAATCA 1020 CAAGTCGAAC CGCCAGAGGT GTAGCTGTCC CTCATGGTAA CCATTACCAC TTTATCCCTT 1080 ATGAACAAT GTCTGAATTG GAAAAACGAA TTGCTCGTAT TATTCCCCTT CGTTATCGTT 1140 CAAACCATTG GGTACCAGAT TCAAGACCAG AACAACCAAG TCCACAATCG ACTCCGGAAC 1200 CTAGTCCAAG TCTGCAACCT GCACCAAATC CTCAACCAGC TCCAAGCAAT CCAATTGATG AGAAATTGGT CAAAGAAGCT GTTCGAAAAG TAGGCGATGG TTATGTCTTT GAGGAGAATG 1320 GAGTTTCTCG TTATATCCCA GCCAAGGATC TTTCAGCAGA AACAGCAGCA GGCATTGATA 1380 GCAAACTGGC CAAGCAGGAA AGTTTATCTC ATAAGCTAGG AGCTAAGAAA ACTGACCTCC 1440 CATCTAGTGA TCGAGAATTT TACAATAAGG CTTATGACTT ACTAGCAAGA ATTCACCAAG 1500 ATTTACTTGA TAATAAAGGT CGACAAGTTG ATTTTGAGGT TTTGGATAAC CTGTTGGAAC 1560 GACTCAAGGA TGTCTCAAGT GATAAAGTCA AGTTAGTGGA TGATATTCTT GCCTTCTTAG 1620 CTCCGATTCG TCATCCAGAA CGTTTAGGAA AACCAAATGC GCAAATTACC TACACTGATG ATGAGATTCA AGTAGCCAAG TTGGCAGGCA AGTACACAAC AGAAGACGGT TATATCTTTG 1740 ATCCTCGTGA TATAACCAGT GATGAGGGGG ATGCCTATGT AACTCCACAT ATGACCCATA 1800 GCCACTGGAT TAAAAAAGAT AGTTTGTCTG AAGCTGAGAG AGCGGCAGCC CAGGCTTATG 1860 CTAAAGAGAA AGGTTTGACC CCTCCTTCGA CAGACCACCA GGATTCAGGA AATACTGAGG 1920 CAAAAGGAGC AGAAGCTATC TACAACCGCG TGAAAGCAGC TAAGAAGGTG CCACTTGATC 1980 GTATGCCTTA CAATCTTCAA TATACTGTAG AAGTCAAAAA CGGTAGTTTA ATCATACCTC 2040 ATTATGACCA TTACCATAAC ATCAAATTTG AGTGGTTTGA CGAAGGCCTT TATGAGGCAC CTAAGGGGTA TAGTCTTGAG GATCTTTTGG CGACTGTCAA GTACTATGTC GAACATCCAA 2160 ACGAACGTCC GCATTCAGAT AATGGTTTTG GTAACGCTAG TGACCATGTT CGTAAAAATA 2220 AGGCAGACCA AGATAGTAAA CCTGATGAAG ATAAGGAACA TGATGAAGTA AGTGAGCCAA 2280 CTCACCCTGA ATCTGATGAA AAAGAGAATC ACGCTGGTTT AAATCCTTCA GCAGATAATC 2340 TTTATAAACC AAGCACTGAT ACGGAAGAGA CAGAGGAAGA AGCTGAAGAT ACCACAGATG 2400 AGGCTGAAAT TCCTCAAGTA GAGAATTCTG TTATTAACGC TAAGATAGCA GATGCGGAGG 2460 CCTTGCTAGA AAAAGTAACA GATCCTAGTA TTAGACAAAA TGCTATGGAG ACATTGACTG GTCTAAAAAG TAGTCTTCTT CTCGGAACGA AAGATAATAA CACTATTTCA GCAGAAGTAG 2580 ATAGTCTCTT GGCTTTGTTA AAAGAAAGTC AACCGGCTCC TATACAGTAG TAAAATGAA 2639 (SEQ ID NO : 13)

FIGURE 16

MKINKKYLAG	SVAVLALSVC	SYELGRHQAG	QVKKESNRVS	YIDGDQAGQK	50
AENLTPDEVS	KREGINAEQI	VIKITDQGYV	TSHGDHYHYY	NGKVPYDAII	100
SEELLMKDPN	YQLKDSDIVN	EIKGGYVIKV	DGKYYVYLKD	AAHADNIRTK	150
EEIKRQKQEH	SHNHNSRADN	AVAAARAQGR	YTTDDGYIFN	ASDIIEDTGD	200
AYIVPHGDHY	HYIPKNELSA	SELAAAEAYW	NGKQGSRPSS	SSSYNANPVQ	250
PRLSENHNLT	VTPTYHQNQG	ENISSLLREL	YAKPLSERHV	ESDGLIFDPA	300
QITSRTARGV	AVPHGNHYHF	IPYEQMSELE	KRIARIIPLR	YRSNHWVPDS	350
RPEQPSPQST	PEPSPSLQPA	PNPQPAPSNP	IDEKLVKEAV	RKVGDGYVFE	400
ENGVSRYIPA	KDLSAETAAG	IDSKLAKQES	LSHKLGAKKT	DLPSSDREFY	450
NKAYDLLARI	HQDLLDNKGR	QVDFEVLDNL	LERLKDVSSD	KVKLVDDILA	500
FLAPIRHPER	LGKPNAQITY	TDDEIQVAKL	AGKYTTEDGY	IFDPRDITSD	550
EGDAYVTPHM	THSHWIKKDS	LSEAERAAAQ	AYAKEKGLTP	PSTDHQDSGN	600
TEAKGAEAIY	NRVKAAKKVP	LDRMPYNLQY	TVEVKNGSLI	IPHYDHYHNI	650
KFEWFDEGLY	EAPKGYSLED	LLATVKYYVE	HPNERPHSDN	GFGNASDHVR	700
KNKADQDSKP	DEDKEHDEVS	EPTHPESDEK	ENHAGLNPSA	DNLYKPSTDT	750
EETEEEAEDT	TDEAEIPQVE	NSVINAKIAD	AEALLEKVTD	PSIRQNAMET	800
LTGLKSSLLL	GTKDNNTISA	EVDSLLALLK	ESQPAPIQ		838
(SEQ ID NO	: 14)				

FIGURE 17

TGTGCCTATG CACTAAACCA GCATCGTTCG CAGGAAAATA AGGACAATAA TCGTGTCTCT 60 TATGTGGATG GCAGCCAGTC AAGTCAGAAA AGTGAAAACT TGACACCAGA CCAGGTTAGC 120 CAGAAAGAAG GAATTCAGGC TGAGCAAATT GTAATCAAAA TTACAGATCA GGGCTATGTA 180 ACGTCACACG GTGATCACTA TCATTACTAT AATGGGAAAG TTCCTTATGA TGCCCTCTTT 240 AGTGAAGAAC TCTTGATGAA GGATCCAAAC TATCAACTTA AAGACGCTGA TATTGTCAAT 300 GAAGTCAAGG GTGGTTATAT CATCAAGGTC GATGGAAAAT ATTATGTCTA CCTGAAAGAT 360 GCAGCTCATG CTGATAATGT TCGAACTAAA GATGAAATCA ATCGTCAAAA ACAAGAACAT 420 GTCAAAGATA ATGAGAAGGT TAACTCTAAT GTTGCTGTAG CAAGGTCTCA GGGACGATAT 480 ACGACAAATG ATGGTTATGT CTTTAATCCA GCTGATATTA TCGAAGATAC GGGTAATGCT 540 TATATCGTTC CTCATGGAGG TCACTATCAC TACATTCCCA AAAGCGATTT ATCTGCTAGT GAATTAGCAG CAGCTAAAGC ACATCTGGCT GGAAAAAATA TGCAACCGAG TCAGTTAAGC 660 TATTCTTCAA CACCTTCTCC ATCTCTTCCA ATCAATCCAG GAACTTCACA TGAGAAACAT 720 GAAGAAGATG GATACGGATT TGATGCTAAT CGTATTATCG CTGAAGATGA ATCAGGTTTT 780 GTCATGAGTC ACGGAGACCA CAATCATTAT TTCTTCAAGA AGGACTTGAC AGAAGAGCAA 840 ATTAAGGCTG CGCAAAAACA TTTAGAGGAA GTTAAAACTA GTCATAATGG ATTAGATTCT 900 TTGTCATCTC ATGAACAGGA TTATCCAAGT AATGCCAAAG AAATGAAAAGA TTTAGATAAA 960 AAAATCGAAG AAAAAATTGC TGGCATTATG AAACAATATG GTGTCAAACG TGAAAGTATT 1020 GTCGTGAATA AAGAAAAAA TGCGATTATT TATCCGCATG GAGATCACCA TCATGCAGAT 1080 CCGATTGATG AACATAAACC GGTTGGAATT GGTCATTCTC ACAGTAACTA TGAACTGTTT 1140 AAACCCGAAG AAGGAGTTGC TAAAAAAGAA GGGAATAAAG TTTATACTGG AGAAGAATTA 1200 ACGAATGTTG TTAATTTGTT AAAAAATAGT ACGTTTAATA ATCAAAACTT TACTCTAGCC 1260 AATGGTCAAA AACGCGTTTC TTTTAGTTTT CCGCCTGAAT TGGAGAAAAA ATTAGGTATC 1320 AATATGCTAG TAAAATTAAT AACACCAGAT GGAAAAGTAT TGGAGAAAGT ATCTGGTAAA GTATTTGGAG AAGGAGTAGG GAATATTGCA AACTTTGAAT TAGATCAACC TTATTTACCA 1440 GGACAACAT TTAAGTATAC TATCGCTTCA AAAGATTATC CAGAAGTAAG TTATGATGGT 1500 ACATTTACAG TTCCAACCTC TTTAGCTTAC AAAATGGCCA GTCAAACGAT TTTCTATCCT 1560 TTCCATGCAG GGGATACTTA TTTAAGAGTG AACCCTCAAT TTGCAGTGCC TAAAGGAACT 1620 GATGCTTTAG TCAGAGTGTT TGATGAATTT CATGGAAATG CTTATTTAGA AAATAACTAT 1680 AAAGTTGGTG AAATCAAATT ACCGATTCCG AAATTAAACC AAGGAACAAC CAGAACGGCC 1740 GGAAATAAAA TTCCTGTAAC CTTCATGGCA AATGCTTATT TGGACAATCA ATCGACTTAT 1860 ATTGTGGAAG TACCTATCTT GGAAAAAGAA AATCAAACTG ATAAACCAAG TATTCTACCA CAATTTAAAA GGAATAAAGC ACAAGAAAAC TCAAAACTTG ATGAAAAGGT AGAAGAACCA 1920 AAGACTAGTG AGAAGGTAGA AAAAGAAAAA CTTTCTGAAA CTGGGAATAG TACTAGTAAT 1980 TCAACGTTAG AAGAAGTTCC TACAGTGGAT CCTGTACAAG AAAAAGTAGC AAAATTTGCT 2040 GAAAGTTATG GGATGAAGCT AGAAAATGTC TTGTTTAATA TGGACGGAAC AATTGAATTA 2100 TATTTACCAT CGGGAGAAGT CATTAAAAAG AATATGGCAG ATTTTACAGG AGAAGCACCT 2160 CAAGGAAATG GTGAAAATAA ACCATCTGAA AATGGAAAAG TATCTACTGG AACAGTTGAG 2220 AACCAACCAA CAGAAAATAA ACCAGCAGAT TCTTTACCAG AGGCACCAAA CGAAAAACCT 2280 GTAAAACCAG AAAACTCAAC GGATAATGGA ATGTTGAATC CAGAAGGGAA TGTGGGGAGT 2340 GACCCTATGT TAGATTCAGC ATTAGAGGAA GCTCCAGCAG TAGATCCTGT ACAAGAAAAA TTAGAAAAAT TTACAGCTAG TTACGGATTA GGCTTAGATA GTGTTATATT CAATATGGAT 2460 GGAACGATTG AATTAAGATT GCCAAGTGGA GAAGTGATAA AAAAGAATTT ATTGATCTCA 2520 (SEQ ID NO : 15) 2528 TAGCGTAA

VIKITDQGYV TSHGDHYHYY NGKVPYDALF SEELLMKDPN YQLKDADIVN	100
EVKGGYIIKV DGKYYVYLKD AAHADNVRTK DEINRQKQEH VKDNEKVNSN	150
VAVARSQGRY TTNDGYVFNP ADIIEDTGNA YIVPHGGHYH YIPKSDLSAS	200
ELAAAKAHLA GKNMQPSQLS YSSTPSPSLP INPGTSHEKH EEDGYGFDAN	250
RIIAEDESGF VMSHGDHNHY FFKKDLTEEQ IKAAQKHLEE VKTSHNGLDS	300
LSSHEQDYPS NAKEMKDLDK KIEEKIAGIM KQYGVKRESI VVNKEKNAII	350
YPHGDHHHAD PIDEHKPVGI GHSHSNYELF KPEEGVAKKE GNKVYTGEEL	400
TNVVNLLKNS TFNNQNFTLA NGQKRVSFSF PPELEKKLGI NMLVKLITPD	450
GKVLEKVSGK VFGEGVGNIA NFELDQPYLP GQTFKYTIAS KDYPEVSYDG	500
TFTVPTSLAY KMASQTIFYP FHAGDTYLRV NPQFAVPKGT DALVRVFDEF	550
HGNAYLENNY KVGEIKLPIP KLNQGTTRTA GNKIPVTFMA NAYLDNQSTY	600
IVEVPILEKE NQTDKPSILP QFKRNKAQEN SKLDEKVEEP KTSEKVEKEK	650
LSETGNSTSN STLEEVPTVD PVQEKVAKFA ESYGMKLENV LFNMDGTIEL	700
YLPSGEVIKK NMADFTGEAP QGNGENKPSE NGKVSTGTVE NQPTENKPAD	750
SLPEAPNEKP VKPENSTDNG MLNPEGNVGS DPMLDSALEE APAVDPVQEK	800
LEKFTASYGL GLDSVIFNMD GTIELRLPSG EVIKKNLLIS	840
(SEQ ID NO : 16)	

CAYALNQHRS	QENKDNNRVS	YVDGSQSSQK	SENLTPDQVS	QKEGIQAEQI	50
VIKITDQGYV	TSHGDHYHYY	NGKVPYDALF	SEELLMKDPN	YQLKDADIVN	100
EVKGGYIIKV	DGKYYVYLKD	AAHADNVRTK	DEINRQKQEH	VKDNEKVNSN	150
VAVARSQGRY	TTNDGYVFNP	ADIIEDTGNA	YIVPHGGHYH	YIPKSDLSAS	200
ELAAAKAHLA	GKNMQPSQLS	YSSTASDNNT	QSVAKGSTSK	PANKSENLQS	250
LLKELYDSPS	AQRYSESDGL	VFDPAKIISR	TPNGVAIPHG	DHYHFIPYSK	300
LSALEEKIAR	MVPISGTGST	VSTNAKPNEV	VSSLGSLSSN	PSSLTTSKEL	350
SSASDGYIFN	PKDIVEETAT	AYIVRHGDHF	HYIPKSNQIG	QPTLPNNSLA	400
TPSPSLPINP	GTSHEKHEED	GYGFDANRII	AEDESGFVMS	HGDHNHYFFK	450
KDLTEEQIKA	AQKHLEEVKT	SHNGLDSLSS	HEQDYPGNAK	EMKDLDKKIE	500
EKIAGIMKQY	GVKRESIVVN	KEKNAIIYPH	GDHHHADPID	EHKPVGIGHS	550
HSNYELFKPE	EGVAKKEGNK	VYTGEELTNV	VNLLKNSTFN	NQNFTLANGQ	600
KRVSFSFPPE	LEKKLGINML	VKLITPDGKV	LEKVSGKVFG	EGVGNIANFE	650
LDQPYLPGQT	FKYTIASKDY	PEVSYDGTFT	VPTSLAYKMA	SQTIFYPFHA	700
GDTYLRVNPQ	FAVPKGTDAL	VRVFDEFHGN	AYLENNYKVG	EIKLPIPKLN	750
QGTTRTAGNK	IPVTFMANAY	LDNQSTYIVE	VPILEKENQT	DKPSILPQFK	800
RNKAQENSKL	DEKVEEPKTS	EKVEKEKLSE	TGNSTSNSTL	EEVPTVDPVQ	850
EKVAKFAESY	GMKLENVLFN	MDGTIELYLP	SGEVIKKNMA	DFTGEAPQGN	900
GENKPSENGK	VSTGTVENQP	TENKPADSLP	EAPNEKPVKP	ENSTDNGMLN	950
PEGNVGSDPM	LDPALEEAPA	VDPVQEKLEK	FTASYGLGLD	SVIFNMDGTI	1000
ELRLPSGEVI	KKNLSDFIA	(SEQ ID NO) : 55)		1019

FIGURE 20

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CAYALNQHRS	QENKDNNRVS	YVDGSQSSQK	SENLTPDQVS	QKEGIQAEQI	50
VIKITDQGYV	TSHGDHYHYY	NGKVPYDALF	SEELLMKDPN	YQLKDADIVN	100
EVKGGYIIKV	DGKYYVYLKD	AAHADNVRTK	DEINRQKQEH	VKDNEKVNSN	150
VAVARSQGRY	TTNDGYVFNP	ADIIEDTGNA	YIVPHGGHYH	YIPKSDLSAS	200
ELAAAKAHLA	${\tt GKNMQPSQLS}$	YSSTASDNNT	QSVAKGSTSK	PANKSENLQS	250
LLKELYDSPS	AQRYSESDGL	VFDPAKIISR	TPNGVAIPHG	DHYHFIPYSK	300
LSALEEKIAR	MVPISGTGST	VSTNAKPNEV	VSSLGSLSSN	PSSLTTSKEL	350
SSASDGYIFN	PKDIVEETAT	AYIVRHGDHF	HYIPKSNQIG	QPTLPNNSLA	400
TPSPSLPINP	GTSHEKHEED	GYGFDANRII	AEDESGFVMS	HGDHNHYFFK	450
KDLTEEQIKA	AQKHLEEVKT	SHNGLDSLSS	HEQDYPGNA		489
(SEQ ID NO	: 56)				

${\tt MKFSKKYIAA}$	GSAVIVSLSL	CAYALNQHRS	QENKDNNRVS	YVDGSQSSQK	SENLTPDQVS	60
QKEGIQAEQI	VIKITDQGYV	TSHGDHYHYY	NGKVPYDALF	SEELLMKDPN	YQLKDADIVN	120
EVKGGYIIKV	$\mathtt{DGKYYVYLKD}$	$\mathtt{AAHADNVRTK}$	DEINRQKQEH	VKDNEKVNSN	VAVARSQGRY	180
TTNDGYVFNP	ADIIEDTGNA	YIVPHGGHYH	YIPKSDLSAS	ELAAAKAHLA	GKNMQPSQLS	240
YSSTASDNNT	QSVAKGSTSK	PANKSENLQS	LLKELYDSPS	AQRYSESDGL	VFDPAKIISR	300
TPNGVAIPHG	DHYHFIPYSK	LSALEEKIAR	MVPISGTGST	VSTNAKPNEV	VSSLGSLSSN	360
PSSLTTSKEL	${\tt SSASDGYIFN}$	PKDIVEETAT	AYIVRHGDHF	HYIPKSNQIG	QPTLPNNSLA	420
TPSPSLPINP	GTSHEKHEED	GYGFDANRII	AEDESGFVMS	${\tt HGDHNHYFFK}$	KDLTEEQIKA	480
AQKHLEEVKT	SHNGLDSLSS	HEQDYPGNA	(SEQ ID NO	o: 57)		509

DLTEEQIKAA	QKHLEEVKTS	HNGLDSLSSH	EQDYPGNAKE	MKDLDKKIEE	50
KIAGIMKQYG	VKRESIVVNK	EKNAIIYPHG	DHHHADPIDE	HKPVGIGHSH	100
SNYELFKPEE	GVAKKEGNKV	YTGEELTNVV	NLLKNSTFNN	QNFTLANGQK	150
RVSFSFPPEL	EKKLGINMLV	KLITPDGKVL	EKVSGKVFGE	GVGNIANFEL	200
DQPYLPGQTF	KYTIASKDYP	EVSYDGTFTV	PTSLAYKMAS	QTIFYPFHAG	250
DTYLRVNPQF	${\tt AVPKGTDALV}$	RVFDEFHGNA	YLENNYKVGE	IKLPIPKLNQ	300
GTTRTAGNKI	PVTFMANAYL	DNQSTYIVEV	PILEKENQTD	KPSILPQFKR	350
NKAQENSKLD	EKVEEPKTSE	KVEKEKLSET	GNSTSNSTLE	EVPTVDPVQE	400
KVAKFAESYG	${\tt MKLENVLFNM}$	DGTIELYLPS	GEVIKKNMAD	FTGEAPQGNG	450
ENKPSENGKV	STGTVENQPT	ENKPADSLPE	${\tt APNEKPVKPE}$	NSTDNGMLNP	500
EGNVGSDPML	DPALEEAPAV	DPVQEKLEKF	TASYGLGLDS	VIFNMDGTIE	550
LRLPSGEVIK	KNLSDFIAKL	RYRSNHWVPD	SRPEEPSPQP	TPEPSPSPQP	600
APNPQPAPSN	PIDEKLVKEA	VRKVGDGYVF	EENGVSRYIP	AKNLSAETAA	650
GIDSKLAKQE	SLSHKLGAKK	TDLPSSDREF	YNKAYDLLAR	IHQDLLDNKG	700
RQVDFEALDN	LLERLKDVSS	DKVKLVDDIL	AFLAPIRHPE	RLGKPNAQIT	750
YTDDEIQVAK	LAGKYTTEDG	YIFDPRDITS	DEGDAYVTPH	MTHSHWIKKD	800
SLSEAERAAA	QAYAKEKGLT	PPSTDHQDSG	NTEAKGAEAI	YNRVKAAKKV	850
PLDRMPYNLQ	YTVEVKNGSL	IIPHYDHYHN	IKFEWFDEGL	YEAPKGYTLE	900
DLLATVKYYV	EHPNERPHSD	NGFGNASDHV	QRNKNGQADT	NQTEKPSEEK	950
PQTEKPEEET	PREEKPQSEK	PESPKPTEEP	EEESPEESEE	PQVETEKVEE	1000
KLREAEDLLG	KIQDPIIKSN	AKETLTGLKN	${\tt NLLFGTQDNN}$	TIMAEAEKLL	1050
ALLKESK	(SEQ ID NO :	: 58)			1057

FIGURE 23

CAYALNOHRS	OENKDNNRVS	YVDGSOSSOK	SENLTPDQVS	OKEGIOAEOI	50
			SEELLMKDPN		100
-			DEINRQKQEH	·-	150
			YIVPHGGHYH		200
	EQ ID NO :	59)			205
(,	-~	,	FIGURE 2	4	
CAYELGLHQA	QTVKENNRVS	YIDGKQATQK	TENLTPDEVS	KREGINAEQI	50
VIKITDQGYV	TSHGDHYHYY	NGKVPYDAII	SEELLMKDPN	YQLKDSDIVN	100
EIKGGYVIKV	NGKYYVYLKD	AAHADNVRTK	EEINRQKQEH	SQHREGGTSA	150
NDGAVAFARS	QGRYTTDDGY	IFNASDIIED	TGDAYIVPHG	DHYHYIPKNE	200
LSASELAAAE	AFLSGRENLS	NLRTYRRQNS	DNTPRTNWVP	SVSNPGTTNT	250
NTSNNSNTNS	QASQSNDIDS	LLKQLYKLPL	SQRHVESDGL	IFDPAQITSR	300
TARGVAVPHG	NHYHFIPYEQ	MSELEKRIAR	IIPLRYRSNH	WVPDSRPEEP	350
SPQPTPEPSP	SPQPAPNPQP	APSNPIDEKL	VKEAVRKVGD	GYVFEENGVS	400
RYIPAKNLSA	ETAAGIDSKL	AKQESLSHKL	GAKKTDLPSS	DREFYNKAYD	450
LLARIHQDLL	DNKGRQVDFE	ALDNLLERLK	DVSSDKVKLV	DDILAFLAPI	500
RHPERLGKPN	AQITYTDDEI	QVAKLAGKYT	TEDGYIFDPR	DITSDEGDAY	550
VTPHMTHSHW	IKKDSLSEAE	RAAAQAYAKE	KGLTPPSTDH	QDSGNTEAKG	600
AEAIYNRVKA	AKKVPLDRMP	YNLQYTVEVK	NGSLIIPHYD	HYHNIKFEWF	650
DEGLYEAPKG	YTLEDLLATV		PHSDNGFGNA		700
OADTNOTEKP	SEEKPOTEKP	EEETPREEKP	QSEKPESPKP	TEEPEEESPE	750
ESEEPQVETE	KVEEKLREAE	DLLGKIQDPI	IKSNAKETLT	GLKNNLLFGT	800
			ID NO : 60)		821
	EKLLALLKES				821
			ID NO : 60)		821
			ID NO : 60)		821
			ID NO : 60)		
QDNNTIMAEA CAYELGLHQA	EKLLALLKES QTVKENNRVS	K ((SEQ :	ID NO : 60) FIGURE 2 TENLTPDEVS	5 KREGINAEQI	50
QDNNTIMAEA CAYELGLHQA VIKITDQGYV	EKLLALLKES QTVKENNRVS TSHGDHYHYY	K ((SEQ T	ID NO : 60) FIGURE 2 TENLTPDEVS SEELLMKDPN	5 KREGINAEQI YQLKDSDIVN	50 100
QDNNTIMAEA CAYELGLHQA VIKITDQGYV EIKGGYVIKV	EKLLALLKES QTVKENNRVS TSHGDHYHYY NGKYYVYLKD	K ((SEQ TO	ID NO : 60) FIGURE 2 TENLTPDEVS SEELLMKDPN EEINRQKQEH	5 KREGINAEQI YQLKDSDIVN SQHREGGTSA	50 100 150
QDNNTIMAEA CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY	K ((SEQ TO SERVICE OF	TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE	50 100 150 200
QDNNTIMAEA CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS	FIGURE 2 FIGURE 2 TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT	50 100 150 200 250
QDNNTIMAEA CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL	FIGURE 2 TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT	50 100 150 200 250 300
QDNNTIMAEA CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS TARGVAVPHG	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS NHYHFIPYEQ	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL	FIGURE 2 TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT	50 100 150 200 250
QDNNTIMAEA CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS NHYHFIPYEQ	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL	FIGURE 2 TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT	50 100 150 200 250 300
QDNNTIMAEA CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS TARGVAVPHG	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS NHYHFIPYEQ	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL	FIGURE 2 TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT IFDPAQITSR	50 100 150 200 250 300
QDNNTIMAEA CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS TARGVAVPHG	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS NHYHFIPYEQ	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL	TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT IFDPAQITSR	50 100 150 200 250 300
QDNNTIMAEA CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS TARGVAVPHG	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS NHYHFIPYEQ	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL	TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT IFDPAQITSR	50 100 150 200 250 300
QDNNTIMAEA CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS TARGVAVPHG (SEQ ID NO	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS NHYHFIPYEQ : 61)	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL MSELEKRIAR	TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL IIPL FIGURE 2	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT IFDPAQITSR	50 100 150 200 250 300 334
QDNNTIMAEA CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS TARGVAVPHG (SEQ ID NO	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS NHYHFIPYEQ : 61)	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL MSELEKRIAR	TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL IIPL FIGURE 2	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT IFDPAQITSR	50 100 150 200 250 300 334
CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS TARGVAVPHG (SEQ ID NO RYRSNHWVPD VRKVGDGYVF	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS NHYHFIPYEQ : 61) SRPEEPSPQP EENGVSRYIP	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL MSELEKRIAR TPEPSPSPQP AKNLSAETAA	TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL IIPL FIGURE 2 APNPQPAPSN GIDSKLAKQE	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT IFDPAQITSR 6 PIDEKLVKEA SLSHKLGAKK	50 100 150 200 250 300 334
CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS TARGVAVPHG (SEQ ID NO RYRSNHWVPD VRKVGDGYVF TDLPSSDREF	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS NHYHFIPYEQ : 61) SRPEEPSPQP EENGVSRYIP YNKAYDLLAR	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL MSELEKRIAR TPEPSPSPQP AKNLSAETAA IHQDLLDNKG	TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL IIPL FIGURE 2 APNPQPAPSN GIDSKLAKQE RQVDFEALDN	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT IFDPAQITSR 6 PIDEKLVKEA SLSHKLGAKK LLERLKDVSS	50 100 150 200 250 300 334
CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS TARGVAVPHG (SEQ ID NO RYRSNHWVPD VRKVGDGYVF TDLPSSDREF DKVKLVDDIL	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS NHYHFIPYEQ : 61) SRPEEPSPQP EENGVSRYIP YNKAYDLLAR AFLAPIRHPE	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL MSELEKRIAR TPEPSPSPQP AKNLSAETAA IHQDLLDNKG RLGKPNAQIT	TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL IIPL FIGURE 2 APNPQPAPSN GIDSKLAKQE RQVDFEALDN YTDDEIQVAK	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT IFDPAQITSR 6 PIDEKLVKEA SLSHKLGAKK LLERLKDVSS LAGKYTTEDG	50 100 150 200 250 300 334 50 100 150 200
CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS TARGVAVPHG (SEQ ID NO RYRSNHWVPD VRKVGDGYVF TDLPSSDREF DKVKLVDDIL YIFDPRDITS	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS NHYHFIPYEQ : 61) SRPEEPSPQP EENGVSRYIP YNKAYDLLAR AFLAPIRHPE DEGDAYVTPH	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL MSELEKRIAR TPEPSPSPQP AKNLSAETAA IHQDLLDNKG RLGKPNAQIT MTHSHWIKKD	TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL IIPL FIGURE 2 APNPQPAPSN GIDSKLAKQE RQVDFEALDN YTDDEIQVAK SLSEAERAAA	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT IFDPAQITSR 6 PIDEKLVKEA SLSHKLGAKK LLERLKDVSS LAGKYTTEDG QAYAKEKGLT	50 100 150 200 250 300 334
CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS TARGVAVPHG (SEQ ID NO RYRSNHWVPD VRKVGDGYVF TDLPSSDREF DKVKLVDDIL YIFDPRDITS PPSTDHQDSG	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS NHYHFIPYEQ : 61) SRPEEPSPQP EENGVSRYIP YNKAYDLLAR AFLAPIRHPE DEGDAYVTPH NTEAKGAEAI	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL MSELEKRIAR TPEPSPSPQP AKNLSAETAA IHQDLLDNKG RLGKPNAQIT MTHSHWIKKD YNRVKAAKKV	TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL IIPL FIGURE 2 APNPQPAPSN GIDSKLAKQE RQVDFEALDN YTDDEIQVAK SLSEAERAAA PLDRMPYNLQ	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT IFDPAQITSR 6 PIDEKLVKEA SLSHKLGAKK LLERLKDVSS LAGKYTTEDG QAYAKEKGLT YTVEVKNGSL	50 100 150 200 250 300 334 50 100 150 200 250 300
CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS TARGVAVPHG (SEQ ID NO RYRSNHWVPD VRKVGDGYVF TDLPSSDREF DKVKLVDDIL YIFDPRDITS PPSTDHQDSG IIPHYDHYHN	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS NHYHFIPYEQ : 61) SRPEEPSPQP EENGVSRYIP YNKAYDLLAR AFLAPIRHPE DEGDAYVTPH NTEAKGAEAI IKFEWFDEGL	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL MSELEKRIAR TPEPSPSPQP AKNLSAETAA IHQDLLDNKG RLGKPNAQIT MTHSHWIKKD YNRVKAAKKV YEAPKGYTLE	TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL IIPL FIGURE 2 APNPQPAPSN GIDSKLAKQE RQVDFEALDN YTDDEIQVAK SLSEAERAAA PLDRMPYNLQ DLLATVKYYV	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT IFDPAQITSR 6 PIDEKLVKEA SLSHKLGAKK LLERLKDVSS LAGKYTTEDG QAYAKEKGLT YTVEVKNGSL EHPNERPHSD	50 100 150 200 250 300 334 50 100 150 200 250 300 350
CAYELGLHQA VIKITDQGYV EIKGGYVIKV NDGAVAFARS LSASELAAAE NTSNNSNTNS TARGVAVPHG (SEQ ID NO RYRSNHWVPD VRKVGDGYVF TDLPSSDREF DKVKLVDDIL YIFDPRDITS PPSTDHQDSG IIPHYDHYHN NGFGNASDHV	QTVKENNRVS TSHGDHYHYY NGKYYVYLKD QGRYTTDDGY AFLSGRENLS QASQSNDIDS NHYHFIPYEQ : 61) SRPEEPSPQP EENGVSRYIP YNKAYDLLAR AFLAPIRHPE DEGDAYVTPH NTEAKGAEAI IKFEWFDEGL QRNKNGQADT	YIDGKQATQK NGKVPYDAII AAHADNVRTK IFNASDIIED NLRTYRRQNS LLKQLYKLPL MSELEKRIAR TPEPSPSPQP AKNLSAETAA IHQDLLDNKG RLGKPNAQIT MTHSHWIKKD YNRVKAAKKV YEAPKGYTLE NQTEKPSEEK	TENLTPDEVS SEELLMKDPN EEINRQKQEH TGDAYIVPHG DNTPRTNWVP SQRHVESDGL IIPL FIGURE 2 APNPQPAPSN GIDSKLAKQE RQVDFEALDN YTDDEIQVAK SLSEAERAAA PLDRMPYNLQ DLLATVKYYV	KREGINAEQI YQLKDSDIVN SQHREGGTSA DHYHYIPKNE SVSNPGTTNT IFDPAQITSR 6 PIDEKLVKEA SLSHKLGAKK LLERLKDVSS LAGKYTTEDG QAYAKEKGLT YTVEVKNGSL EHPNERPHSD PREEKPQSEK	50 100 150 200 250 300 334 50 100 150 200 250 300

AKETLTGLKN NLLFGTQDNN TIMAEAEKLL ALLKESK

(SEQ ID NO : 62)

487

AEAFLSGREN	LSNLRTYRRQ	NSDNTPRTNW	VPSVSNPGTT	NTNTSNNSNT	50
		PLSQRHVESD			100
		ARIIPLRYRS			150
SPSPQPAPNP	QPAPSNPIDE	KLVKEAVRKV	GDGYVFEENG	VSRYIPAKNL	200
SAETAAGIDS	KLAKQESLSH	KLGAKKTDLP	SSDREFYNKA	YDLLARIHQD	250
LLDNKGRQVD	FEALDNLLER	LKDVSSDKVK	LVDDILAFLA	PIRHPERLGK	300
PNAQITYTDD	EIQVAKLAGK	YTTEDGYIFD	PRDITSDEGD	AYVTPHMTHS	350
		KEKGLTPPST			400
KAAKKVPLDR	MPYNLQYTVE	VKNGSLIIPH	YDHYHNIKFE	WFDEGLYEAP	450
KGYTLEDLLA	TVKYYVEHPN	ERPHSDNGFG	NASDHVQRNK	NGQADTNQTE	500
KPSEEKPQTE	KPEEETPREE	KPQSEKPESP	KPTEEPEEES	PEESEEPQVE	550
TEKVEEKLRE	AEDLLGKIQD	PIIKSNAKET	LTGLKNNLLF	GTQDNNTIMA	600
EAEKLLALLK		ID NO : 63)			613
			FIGURE 2	8	
DITEROIKAA	OKHLEEVKTS	HNGLDSLSSH	EODYPGNAKE	MKDLDKKIEE	50
		EKNAIIYPHG			100
		YTGEELTNVV			150
		KLITPDGKVL			200
		EVSYDGTFTV			250
		RVFDEFHGNA			300
		DNQSTYIVEV			350
		KVEKEKLSET			400
		DGTIELYLPS			450
		ENKPADSLPE			500
		DPVQEKLEKF			550
	KNLSDFIA				568
DRDI DOD VIR	14,2021 21.	(D=x	FIGURE 2	9	
			1100112 2	-	
DLTEEOIKAA	OKHLEEVKTS	HNGLDSLSSH	EQDYPGNAKE	MKDLDKKIEE	50
		EKNAIIYPHG			100
		YTGEELTNVV			150
		KLITPDGKVL			200
		EVSYDGTFTV			250
		RVFDEFHGNA			300
	PVTFMANAYL		(SEQ ID N		329
TUMONIT	TATTIMATI	714×11	(DEQ ID IN		

(SEQ ID NO : 70)

ETGNSTSNST PSGEVIKKNM PEAPNEKPVK	TDKPSILPQF LEEVPTVDPV ADFTGEAPQG PENSTDNGML DSVIFNMDGT : 66)	QEKVAKFAES NGENKPSENG NPEGNVGSDP	YGMKLENVLF KVSTGTVENQ MLDPALEEAP	NMDGTIELYL PTENKPADSL AVDPVQEKLE	50 100 150 200 240
PYEQMSELEK NPQPAPSNPI DSKLAKQESL VDFEALDNLL DDEIQVAKLA SEAERAAAQA DRMPYNLQYT LATVKYYVEH TEKPEEETPR REAEDLLGKI	KLPLSQRHVE RIARIIPLRY DEKLVKEAVR SHKLGAKKTD ERLKDVSSDK GKYTTEDGYI YAKEKGLTPP VEVKNGSLII PNERPHSDNG EEKPQSEKPE QDPIIKSNAK	RSNHWVPDSR KVGDGYVFEE LPSSDREFYN VKLVDDILAF FDPRDITSDE STDHQDSGNT PHYDHYHNIK FGNASDHVQR SPKPTEEPEE ETLTGLKNNL	PEEPSPQPTP NGVSRYIPAK KAYDLLARIH LAPIRHPERL GDAYVTPHMT EAKGAEAIYN FEWFDEGLYE NKNGQADTNQ ESPEESEEPQ	EPSPSPQPAP NLSAETAAGI QDLLDNKGRQ GKPNAQITYT HSHWIKKDSL RVKAAKKVPL APKGYTLEDL TEKPSEEKPQ VETEKVEEKL MAEAEKLLAL	50 100 150 200 250 300 350 400 450 500 555
PYEQMSELEK NPQPAPSNPI DSKLAKQESL VDFEALDNLL DDEIQVAKLA SEAERAAAQA DRMPYNLQYT	KLPLSQRHVE RIARIIPLRY DEKLVKEAVR SHKLGAKKTD ERLKDVSSDK GKYTTEDGYI YAKEKGLTPP VEVKNGSLII PNERPHSDNG	RSNHWVPDSR KVGDGYVFEE LPSSDREFYN VKLVDDILAF FDPRDITSDE STDHQDSGNT PHYDHYHNIK	PEEPSPQPTP NGVSRYIPAK KAYDLLARIH LAPIRHPERL GDAYVTPHMT EAKGAEAIYN	EPSPSPQPAP NLSAETAAGI QDLLDNKGRQ GKPNAQITYT HSHWIKKDSL RVKAAKKVPL APKGYTLEDL : 68)	50 100 150 200 250 300 350 400 428
DTNQTEKPSE		ETPREEKPQS	EKPESPKPTE	HVQRNKNGQA EPEEESPEES 4	50 100 121
PTEEPEEESP	EESEEPQVET TQDNNTIMAE	EKVEEKLREA	EDLLGKIQDP	PQSEKPESPK IIKSNAKETL	50 100 132

PYEQMSELEK NPQPAPSNPI DSKLAKQESL	KLPLSQRHVE RIARIIPLRY DEKLVKEAVR SHKLGAKKTD ERLKDVSSDK	RSNHWVPDSR KVGDGYVFEE LPSSDREFYN	PEEPSPQPTP NGVSRYIPAK	EPSPSPQPAP NLSAETAAGI QDLLDNKGRQ 71)	50 100 150 200 226
ITSDEGDAYV DSGNTEAKGA YHNIKFEWFD	HPERLGKPNA TPHMTHSHWI EAIYNRVKAA EGLYEAPKGY ID NO : 72)	KKDSLSEAER KKVPLDRMPY TLEDLLATVK	AAAQAYAKEK NLQYTVEVKN	GLTPPSTDHQ GSLIIPHYDH HSDNGFGNAS	50 100 150 200 203
IVIKITDQGY NEIKGGYVIK NAVAAARAQG ASELAAAEAY GENISSLLRE FIPYEQMSEL APNPQPAPSN GIDSKLAKQE RQVDFEVLDN YTDDEIQVAK SLSEAERAAA PLDRMPYNLQ DLLATVKYYV SEPTHPESDE ENSVINAKIA	GQVKKESNRV VTSHGDHYHY VDGKYYVYLK RYTTDDGYIF WNGKQGSRPS LYAKPLSERH EKRIARIIPL PIDEKLVKEA SLSHKLGAKK LLERLKDVSS LAGKYTTEDG QAYAKEKGLT YTVEVKNGSL EHPNERPHSD KENHAGLNPS DAEALLEKVT KESQPAPIQ	YNGKVPYDAI DAAHADNIRT NASDIIEDTG SSSSYNANPV VESDGLIFDP RYRSNHWVPD VRKVGDGYVF TDLPSSDREF DKVKLVDDIL YIFDPRDITS PPSTDHQDSG IIPHYDHYHN NGFGNASDHV ADNLYKPSTD DPSIRQNAME	ISEELLMKDP KEEIKRQKQE DAYIVPHGDH QPRLSENHNL AQITSRTARG SRPEQPSPQS EENGVSRYIP YNKAYDLLAR AFLAPIRHPE DEGDAYVTPH NTEAKGAEAI IKFEWFDEGL RKNKADQDSK TEETEEEAED TLTGLKSSLL	NYQLKDSDIV HSHNHNSRAD YHYIPKNELS TVTPTYHQNQ VAVPHGNHYH TPEPSPSLQP AKDLSAETAA IHQDLLDNKG RLGKPNAQIT MTHSHWIKKD YNRVKAAKKV YEAPKGYSLE PDEDKEHDEV TTDEAEIPQV LGTKDNNTIS	50 100 150 200 250 300 350 400 450 500 650 700 750 800 819
IPYEQMSELE PNPQPAPSNP IDSKLAKQES QVDFEVLDNL TDDEIQVAKL LSEAERAAAQ LDRMPYNLQY LLATVKYYVE EPTHPESDEK	IDEKLVKEAV LSHKLGAKKT LERLKDVSSD AGKYTTEDGY AYAKEKGLTP TVEVKNGSLI HPNERPHSDN ENHAGLNPSA AEALLEKVTD	YRSNHWVPDS RKVGDGYVFE DLPSSDREFY KVKLVDDILA IFDPRDITSD PSTDHQDSGN IPHYDHYHNI GFGNASDHVR DNLYKPSTDT	RPEQPSPQST ENGVSRYIPA NKAYDLLARI FLAPIRHPER EGDAYVTPHM TEAKGAEAIY KFEWFDEGLY KNKADQDSKP EETEEEAEDT LTGLKSSLLL	PEPSPSLQPA KDLSAETAAG HQDLLDNKGR LGKPNAQITY THSHWIKKDS NRVKAAKKVP EAPKGYSLED DEDKEHDEVS TDEAEIPQVE GTKDNNTISA	50 100 150 200 250 300 350 400 450 500 568

VRKNKADQDS KPDEDKEHDE VSEPTHPESD EKENHAGLNP SADNLYKPST 50
DTEETEEEAE DTTDEAEIPQ VENSVINAKI ADAEALLEKV TDPSIRQNAM 100
ETLTGLKSSL LLGTKDNNTI SAEVDSLLAL LKESQPAPIQ 140
(SEQ ID NO : 75)

GACTTGACAG	AAGAGCAAAT	TAAGGCTGCG	CAAAAACATT	TAGAGGAAGT	50
TAAAACTAGT	CATAATGGAT	TAGATTCTTT	GTCATCTCAT	GAACAGGATT	100
ATCCAGGTAA	TGCCAAAGAA	ATGAAAGATT	TAGATAAAAA	AATCGAAGAA	150
AAAATTGCTG	GCATTATGAA	ACAATATGGT	GTCAAACGTG	AAAGTATTGT	200
CGTGAATAAA	GAAAAAAATG	CGATTATTTA	TCCGCATGGA	GATCACCATC	250
ATGCAGATCC	GATTGATGAA	CATAAACCGG	TTGGAATTGG	TCATTCTCAC	300
AGTAACTATG	AACTGTTTAA	ACCCGAAGAA	GGAGTTGCTA	AAAAAGAAGG	350
GAATAAAGTT	TATACTGGAG	AAGAATTAAC	GAATGTTGTT	AATTTGTTAA	400
AAAATAGTAC	GTTTAATAAT	CAAAACTTTA	CTCTAGCCAA	TGGTCAAAAA	450
CGCGTTTCTT	TTAGTTTTCC	GCCTGAATTG	GAGAAAAAAT	TAGGTATCAA	500
TATGCTAGTA	AAATTAATAA	CACCAGATGG	AAAAGTATTG	GAGAAAGTAT	550
CTGGTAAAGT	ATTTGGAGAA	GGAGTAGGGA	ATATTGCAAA	CTTTGAATTA	600
GATCAACCTT	ATTTACCAGG	ACAAACATTT	AAGTATACTA	TCGCTTCAAA	650
AGATTATCCA	GAAGTAAGTT	ATGATGGTAC	ATTTACAGTT	CCAACCTCTT	700
TAGCTTACAA	AATGGCCAGT	CAAACGATTT	TCTATCCTTT	CCATGCAGGG	750
GATACTTATT	TAAGAGTGAA	CCCTCAATTT	GCAGTGCCTA		800
TGCTTTAGTC	AGAGTGTTTG	ATGAATTTCA		TATTTAGAAA	850
ATAACTATAA	AGTTGGTGAA	ATCAAATTAC	CGATTCCGAA	ATTAAACCAA	900
GGAACAACCA	GAACGGCCGG	AAATAAAATT	CCTGTAACCT	TCATGGCAAA	950
TGCTTATTTG	GACAATCAAT	CGACTTATAT	TGTGGAAGTA		1000
AAAAAGAAAA	TCAAACTGAT	AAACCAAGTA	TTCTACCACA	ATTTAAAAGG	1050
AATAAAGCAC	AAGAAAACTC	AAAACTTGAT	GAAAAGGTAG		1100
GACTAGTGAG	AAGGTAGAAA	AAGAAAAACT	TTCTGAAACT	GGGAATAGTA	1150
CTAGTAATTC	AACGTTAGAA	GAAGTTCCTA	CAGTGGATCC	TGTACAAGAA	1200
AAAGTAGCAA	AATTTGCTGA	AAGTTATGGG	ATGAAGCTAG	AAAATGTCTT	1250
	GACGGAACAA				1300
TTAAAAAGAA			AAGCACCTCA		1350
GAAAATAAAC	CATCTGAAAA	TGGAAAAGTA	TCTACTGGAA	CAGTTGAGAA	1400
CCAACCAACA	GAAAATAAAC		TTTACCAGAG		1450
AAAAACCTGT	AAAACCAGAA	AACTCAACGG	ATAATGGAAT	GTTGAATCCA	1500
GAAGGGAATG	TGGGGAGTGA	CCCTATGTTA	GATCCAGCAT	TAGAGGAAGC	1550
TCCAGCAGTA	GATCCTGTAC	AAGAAAAATT	AGAAAAATTT	ACAGCTAGTT	1600
ACGGATTAGG	CTTAGATAGT	GTTATATTCA	ATATGGATGG	AACGATTGAA	1650
TTAAGATTGC	CAAGTGGAGA	AGTGATAAAA	AAGAATTTAT	CTGATTTCAT	1700
AGCGAAGCTT	CGTTATCGTT	CAAACCATTG	GGTACCAGAT	TCAAGACCAG	1750
AAGAACCAAG	TCCACAACCG	ACTCCAGAAC	CTAGTCCAAG	TCCGCAACCT	1800
GCACCAAATC	CTCAACCAGC	TCCAAGCAAT	CCAATTGATG	AGAAATTGGT	1850
CAAAGAAGCT	GTTCGAAAAG	TAGGCGATGG	TTATGTCTTT	GAGGAGAATG	1900
GAGTTTCTCG	TTATATCCCA	GCCAAGAATC	TTTCAGCAGA	AACAGCAGCA	1950
GGCATTGATA	GCAAACTGGC	CAAGCAGGAA	AGTTTATCTC	ATAAGCTAGG	2000
	ACTGACCTCC				2050
	ACTAGCAAGA				2100
CGACAAGTTG	ATTTTGAGGC	TTTGGATAAC	CTGTTGGAAC	GACTCAAGGA	2150
	GATAAAGTCA				2200
	TCATCCAGAA				2250
	ATGAGATTCA				2300
AGAAGACGGT					2350
ATGCCTATGT					2400

Δርጥጥጥርጥርጥር	AAGCTGAGAG	AGCGGCAGCC	СУСССТТУТС	CTAAAGAGAA	2450
AGGTTTTGACC	CCTCCTTCGA	CAGACCATCA	GGATTCAGGA	AATACTGAGG	2500
CAAAAGGAGC	AGAAGCTATC	TACAACCGCG	TGAAAGCAGC	TAAGAAGGTG	2550
CCACTTGATC	GTATGCCTTA	CAATCTTCAA	TATACTGTAG	AAGTCAAAAA	2600
CGGTAGTTTA	ATCATACCTC	ATTATGACCA	TTACCATAAC	ATCAAATTTG	2650
AGTGGTTTGA	CGAAGGCCTT	TATGAGGCAC	CTAAGGGGTA	TACTCTTGAG	2700
GATCTTTTGG	CGACTGTCAA	GTACTATGTC	GAACATCCAA	ACGAACGTCC	2750
GCATTCAGAT	AATGGTTTTG	GTAACGCTAG	CGACCATGTT	CAAAGAAACA	2800
AAAATGGTCA	AGCTGATACC	AATCAAACGG	AAAAACCAAG	CGAGGAGAAA	2850
CCTCAGACAG	AAAAACCTGA	GGAAGAAACC	CCTCGAGAAG	AGAAACCACA	2900
AAGCGAGAAA	CCAGAGTCTC	CAAAACCAAC	AGAGGAACCA	GAAGAAGAAT	2950
CACCAGAGGA	ATCAGAAGAA	CCTCAGGTCG	AGACTGAAAA	GGTTGAAGAA	3000
AAACTGAGAG	AGGCTGAAGA	TTTACTTGGA	AAAATCCAGG	ATCCAATTAT	3050
CAAGTCCAAT	GCCAAAGAGA	CTCTCACAGG	ATTAAAAAAT	AATTTACTAT	3100
TTGGCACCCA	GGACAACAAT	ACTATTATGG	CAGAAGCTGA	AAAACTATTG	3150
GCTTTATTAA	AGGAGAGTAA	G (SEQ II	NO: 76)		3171

EAYWNGKQGS	${\tt RPSSSSSYNA}$	NPVQPRLSEN	${\tt HNLTVTPTYH}$	QNQGENISSL	50
LRELYAKPLS	ERHVESDGLI	FDPAQITSRT	${\tt ARGVAVPHGN}$	HYHFIPYEQM	100
SELEKRIARI	IPLRYRSNHW	VPDSRPEQPS	PQSTPEPSPS	LQPAPNPQPA	150
PSNPIDEKLV	KEAVRKVGDG	YVFEENGVSR	YIPAKDLSAE	TAAGIDSKLA	200
KQESLSHKLG	AKKTDLPSSD	${\tt REFYNKAYDL}$	LARIHQDLLD	NKGRQVDFEV	250
LDNLLERLKD	VSSDKVKLVD	DILAFLAPIR	${\tt HPERLGKPNA}$	QITYTDDEIQ	300
VAKLAGKYTT	EDGYIFDPRD	ITSDEGDAYV	TPHMTHSHWI	KKDSLSEAER	350
AAAQAYAKEK	${\tt GLTPPSTDHQ}$	DSGNTEAKGA	EAIYNRVKAA	KKVPLDRMPY	400
NLQYTVEVKN	${\tt GSLIIPHYDH}$	YHNIKFEWFD	EGLYEAPKGY	SLEDLLATVK	450
YYVEHPNERP	HSDNGFGNAS	DHV (SEQ	ID NO : 77)		473

FIGURE 42

(SEQ ID NO : 79)

CAYALNQHRS QENKDNNRVS YVDGSQSSQK SENLTPDOVS OKEGIOAEOI 50 VIKITDQGYV TSHGDHYHYY NGKVPYDALF SEELLMKDPN YOLKDADIVN 100 EVKGGYIIKV DGKYYVYLKD AAHADNVRTK DEINRQKQEH VKDNEKVNSN 150 VAVARSQGRY TTNDGYVFNP ADIIEDTGNA YIVPHGGHYH YIPKSDLSAS 200 ELAAAKAHLA GKNMQPSQLS YSSTASDNNT QSVAKGSTSK PANKSENLQS 250 LLKELYDSPS AQRYSESDGL VFDPAKIISR TPNGVAIPHG DHYHFIPYSK 300 LSALEEKIAR MVPISGTGST VSTNAKPNEV VSSLGSLSSN PSSLTTSKEL 350 SSASDGYIFN PKDIVEETAT AYIVRHGDHF HYIPKSNQIG QPTLPNNSLA 400 TPSPSLPINP GTSHEKHEED GYGFDANRII AEDESGFVMS HGDHNHYFFK 450 KDLTEEQIKA AQKHLEEVKT SHNGLDSLSS HEQDYPGNAK EMKDLDKKIE 500 EKIAGIMKQY GVKRESIVVN KEKNAIIYPH GDHHHADPID EHKPVGIGHS 550 HSNYELFKPE EGVAKKEGNK VYTGEELTNV VNLLKNSTFN NONFTLANGO 600 KRVSFSFPPE LEKKLGINML VKLITPDGKV LEKVSGKVFG EGVGNIANFE 650 LDQPYLPGQT FKYTIASKDY PEVSYDGTFT VPTSLAYKMA SQTIFYPFHA 700 GDTYLRVNPQ FAVPKGTDAL VRVFDEFHGN AYLENNYKVG EIKLPIPKLN 750 QGTTRTAGNK IPVTFMANAY LDNQSTYIVE (SEQ ID NO : 78) 780 FIGURE 43 CAYELGLHQA QTVKENNRVS YIDGKQATQK TENLTPDEVS KREGINAEQI 50 VIKITDQGYV TSHGDHYHYY NGKVPYDAII SEELLMKDPN YQLKDSDIVN 100 EIKGGYVIKV NGKYYVYLKD AAHADNVRTK EEINRQKQEH SQHREGGTSA NDGAVAFARS QGRYTTDDGY IFNASDIIED TGDAYIVPHG DHYHYIPKNE 200 LSASELAAAE AFLSGRENLS NLRTYRRQNS DNTPRTNWVP SVSNPGTTNT 250 NTSNNSNTNS QASQSNDIDS LLKQLYKLPL SQRHVESDGL IFDPAQITSR 300 TARGVAVPHG NHYHFIPYEO MSELEKRIAR IIPLRYRSNH WVPDSRPEEP 350 SPQPTPEPSP SPQPAPNPQP APSNPIDEKL VKEAVRKVGD GYVFEENGVS 400 RYIPAKNLSA ETAAGIDSKL AKQESLSHKL GAKKTDLPSS DREFYNKAYD 450 LLARIHODLL DNKGROVDFE ALDNLLERLK DVSSDKVKLV DDILAFLAPI 500 RHPERLGKPN AQITYTDDEI OVAKLAGKYT TEDGYIFDPR DITSDEGDAY 550 VTPHMTHSHW IKKDSLSEAE RAAAQAYAKE KGLTPPSTDH QDSGNTEAKG 600 AEAIYNRVKA AKKVPLDRMP YNLQYTVEVK NGSLIIPHYD HYHNIKFEWF 650 DEGLYEAPKG YTLEDLLATV KYYVEHPNER PHSDNGFGNA 690

FIGURE 44

GTGAAGAAA CATATGGTTA TATCGGCTCA GTTGCTGCCA TTTTACTAGC TACTCATATT 60 GGAAGTTACC AACTTGGTAA GCATCATATG GGTCTAGCAA CAAAGGACAA TCAGATTGCC 120 TATATTGATG ACAGCAAAGG TAAGGCAAAA GCCCCTAAAA CAAACAAAAC GATGGATCAA 180 ATCAGTGCTG AAGAAGGCAT CTCTGCTGAA CAGATCGTAG TCAAAATTAC TGACCAAGGC 240 TATGTGACCT CACACGGTGA CCATTATCAT TTTTACAATG GGAAAGTTCC TTATGATGCG 300 ATTATTAGTG AAGAGTTGTT GATGACGGAT CCTAATTACC GTTTTAAACA ATCAGACGTT ATCAATGAAA TCTTAGACGG TTACGTTATT AAAGTCAATG GCAACTATTA TGTTTACCTC 420 AAGCCAGGTA GTAAGCGCAA AAACATTCGA ACCAAACAAC AAATTGCTGA GCAAGTAGCC 480 AAAGGAACTA AAGAAGCTAA AGAAAAAGGT TTAGCTCAAG TGGCCCATCT CAGTAAAGAA 540 GAAGTTGCGG CAGTCAATGA AGCAAAAAGA CAAGGACGCT ATACTACAGA CGATGGCTAT 600 ATTTTTAGTC CGACAGATAT CATTGATGAT TTAGGAGATG CTTATTTAGT ACCTCATGGT 660 AATCACTATC ATTATATTCC TAAAAAGGAT TTGTCTCCAA GTGAGCTAGC TGCTGCACAA 720 GCCTACTGGA GTCAAAAACA AGGTCGAGGT GCTAGACCGT CTGATTACCG CCCGACACCA 780 GCCCCAGGTC GTAGGAAAGC CCCAATTCCT GATGTGACGC CTAACCCTGG ACAAGGTCAT 840 CAGCCAGATA ACGGTGGCTA TCATCCAGCG CCTCCTAGGC CAAATGATGC GTCACAAAAC 900 AAACACCAAA GAGATGAGTT TAAAGGAAAA ACCTTTAAGG AACTTTTAGA TCAACTACAC 960 CGTCTTGATT TGAAATACCG TCATGTGGAA GAAGATGGGT TGATTTTTGA ACCGACTCAA 1020 GTGATCAAAT CAAACGCTTT TGGGTATGTG GTGCCTCATG GAGATCATTA TCATATTATC 1080 CCAAGAAGTC AGTTATCACC TCTTGAAATG GAATTAGCAG ATCGATACTT AGCTGGCCAA 1140 ACTGAGGACA ATGACTCAGG TTCAGAGCAC TCAAAACCAT CAGATAAAGA AGTGACACAT 1200 ACCTTTCTTG GTCATCGCAT CAAAGCTTAC GGAAAAGGCT TAGATGGTAA ACCATATGAT 1260 ACGAGTGATG CTTATGTTTT TAGTAAAGAA TCCATTCATT CAGTGGATAA ATCAGGAGTT 1320 ACAGCTAAAC ACGGAGATCA TTTCCACTAT ATAGGATTTG GAGAACTTGA ACAATATGAG TTGGATGAGG TCGCTAACTG GGTGAAAGCA AAAGGTCAAG CTGATGAGCT TGCTGCTGCT 1440 TTGGATCAGG AACAAGGCAA AGAAAAACCA CTCTTTGACA CTAAAAAAGT GAGTCGCAAA 1500 GTAACAAAAG ATGGTAAAGT GGGCTATATG ATGCCAAAAG ATGGTAAGGA CTATTTCTAT 1560 GCTCGTGATC AACTTGATTT GACTCAGATT GCCTTTGCCG AACAAGAACT AATGCTTAAA 1620 GATAAGAAGC ATTACCGTTA TGACATTGTT GACACAGGTA TTGAGCCACG ACTTGCTGTA 1680 GATGTGTCAA GTCTGCCGAT GCATGCTGGT AATGCTACTT ACGATACTGG AAGTTCGTTT 1740 GTTATCCCAC ATATTGATCA TATCCATGTC GTTCCGTATT CATGGTTGAC GCGCGATCAG 1800 ATTGCAACAG TCAAGTATGT GATGCAACAC CCCGAAGTTC GTCCGGATGT ATGGTCTAAG 1860 CCAGGGCATG AAGAGTCAGG TTCGGTCATT CCAAATGTTA CGCCTCTTGA TAAACGTGCT 1920 GGTATGCCAA ACTGGCAAAT TATCCATTCT GCTGAAGAAG TTCAAAAAGC CCTAGCAGAA 1980 GGTCGTTTTG CAACACCAGA CGGCTATATT TTCGATCCAC GAGATGTTTT GGCCAAAGAA 2040 ACTTTTGTAT GGAAAGATGG CTCCTTTAGC ATCCCAAGAG CAGATGGCAG TTCATTGAGA 2100 ACCATTAATA AATCTGATCT ATCCCAAGCT GAGTGGCAAC AAGCTCAAGA GTTATTGGCA AAGAAAAATA CTGGTGATGC TACTGATACG GATAAACCCA AAGAAAAGCA ACAGGCAGAT 2220 AAGAGCAATG AAAACCAACA GCCAAGTGAA GCCAGTAAAG AAGAAAAAGA ATCAGATGAC 2280 TTTATAGACA GTTTACCAGA CTATGGTCTA GATAGAGCAA CCCTAGAAGA TCATATCAAT CAATTAGCAC AAAAAGCTAA TATCGATCCT AAGTATCTCA TTTTCCAACC AGAAGGTGTC 2400 CAATTTTATA ATAAAAATGG TGAATTGGTA ACTTATGATA TCAAGACACT TCAACAAATA 2460 AACCCTTAA (SEQ ID NO : 80) 2469

FIGURE 45

VKKTYGYIGS	VAAILLATHI	GSYQLGKHHM	GLATKDNQIA	YIDDSKGKAK	50
APKTNKTMDQ	ISAEEGISAE	QIVVKITDQG	YVTSHGDHYH	FYNGKVPYDA	100
IISEELLMTD	PNYRFKQSDV	INEILDGYVI	KVNGNYYVYL	KPGSKRKNIR	150
TKQQIAEQVA	KGTKEAKEKG	LAQVAHLSKE	EVAAVNEAKR	QGRYTTDDGY	200
IFSPTDIIDD	LGDAYLVPHG	NHYHYIPKKD	LSPSELAAAQ	AYWSQKQGRG	250
ARPSDYRPTP	APGRRKAPIP	DVTPNPGQGH	QPDNGGYHPA	PPRPNDASQN	300
KHQRDEFKGK	TFKELLDQLH	RLDLKYRHVE	EDGLIFEPTQ	VIKSNAFGYV	350
VPHGDHYHII	PRSQLSPLEM	ELADRYLAGQ	TEDNDSGSEH	SKPSDKEVTH	400
TFLGHRIKAY	GKGLDGKPYD	TSDAYVFSKE	SIHSVDKSGV	TAKHGDHFHY	450
IGFGELEQYE	LDEVANWVKA	KGQADELAAA	LDQEQGKEKP	LFDTKKVSRK	500
VTKDGKVGYM	MPKDGKDYFY	ARDQLDLTQI	AFAEQELMLK	DKKHYRYDIV	550
DTGIEPRLAV	DVSSLPMHAG	NATYDTGSSF	VIPHIDHIHV	VPYSWLTRDQ	600
IATVKYVMQH	PEVRPDVWSK	PGHEESGSVI	PNVTPLDKRA	GMPNWQIIHS	650
AEEVQKALAE	GRFATPDGYI	FDPRDVLAKE	TFVWKDGSFS	IPRADGSSLR	700
TINKSDLSQA	EWQQAQELLA	KKNTGDATDT	DKPKEKQQAD	KSNENQQPSE	750
ASKEEKESDD	FIDSLPDYGL	DRATLEDHIN	QLAQKANIDP	KYLIFQPEGV	800
QFYNKNGELV	TYDIKTLQQI	NPP (SEQ	ID NO : 81)		823

FIGURE 46

GTGAAGAAAA CATATGGTTA TATCGGCTCA GTTGCTGCCA TTTTACTAGC TACTCATATT 60 GGAAGTTACC AACTTGGTAA GCATCATATG GGTCTAGCAA CAAAGGACAA TCAGATTGCC 120 TATATTGATG ATAGCAAAGG TAAGGCAAAA GCCCCTAAAA CAAACAAAAC GATGGATCAA 180 ATCAGTGCTG AAGAAGGCAT CTCTGCTGAA CAGATCGTAG TCAAAATTAC TGACCAAGGT 240 TATGTGACCT CACACGGTGA CCATTATCAT TTTTACAATG GGAAAGTTCC TTATGATGCG 300 ATTATTAGTG AAGAGTTGTT GATGACGGAT CCTAATTACC ATTTTAAACA ATCAGACGTT 360 ATCAATGAAA TCTTAGACGG TTACGTTATT AAAGTCAATG GCAACTATTA TGTTTACCTC 420 AAGCCAGGTA GTAAGCGCAA AAACATTCGA ACCAAACAAC AAATTGCTGA GCAAGTAGCC AAAGGAACTA AAGAAGCTAA AGAAAAAGGT TTAGCTCAAG TGGCCCATCT CAGTAAAGAA 540 GAAGTTGCGG CAGTCAATGA AGCAAAAAGA CAAGGACGCT ATACTACAGA CGATGGCTAT 600 ATTTTTAGTC CGACAGATAT CATTGATGAT TTAGGAGACG CTTATTTAGT ACCTCATGGT 660 AATCACTATC ATTATATTCC TAAAAAAGAT TTGTCTCCAA GTGAGCTAGC TGCTGCACAA 720 GCTTACTGGA GTCAAAAACA AGGTCGAGGT GCTAGACCGT CTGATTACCG CCCGACACCA 780 GCCCCAGGTC GTAGGAAAGC TCCAATTCCT GATGTGACGC CTAACCCTGG ACAAGGTCAT 840 CAGCCAGATA ACGGTGGCTA TCATCCAGCG CCTCCTAGGC CAAATGATGC GTCACAAAAC 900 AAACACCAAA GAGATGAGTT TAAAGGAAAA ACCTTTAAGG AACTTTTAGA TCAACTACAC 960 CGTCTTGATT TGAAATACCG TCATGTGGAA GAAGATGGGT TGATTTTTGA ACCGACTCAA 1020 GTGATCAAAT CAAACGCTTT TGGGTATGTG GTGCCTCATG GAGATCATTA TCATATTATC 1080 CCAAGAAGTC AGTTATCACC TCTTGAAATG GAATTAGCAG ATCGATACTT AGCCGGTCAA 1140 ACTGAGGACA ATGATTCAGG TTCAGATCAC TCAAAACCAT CAGATAAAGA AGTGACACAT ACCTTTCTTG GTCATCGCAT CAAAGCTTAC GGAAAAGGCT TAGATGGTAA ACCATATGAT 1260 ACGAGTGATG CTTATGTTTT TAGTAAAGAA TCCATTCATT CAGTGGATAA ATCAGGAGTT 1320 ACAGCTAAAC ACGGAGATCA TTTCCACTAT ATAGGATTTG GAGAACTTGA ACAATATGAG 1380 TTGGATGAGG TCGCTAACTG GGTGAAAGCA AAAGGTCAAG CTGATGAGCT TGCTGCTGCT 1440 TTGGATCAGG AACAAGGCAA AGAAAAACCA CTCTTTGACA CTAAAAAAGT GAGTCGCAAA 1500 GTAACAAAAG ATGGTAAAGT GGGCTATATT ATGCCAAAAG ATGGCAAGGA CTATTTCTAT 1560 GCTCGTGATC AACTTGATTT GACTCAGATT GCCTTTGCCG AACAAGAACT AATGCTTAAA GATAAGAACC ATTACCGTTA TGACATTGTT GACACAGGTA TTGAGCCACG ACTTGCTGTA 1680 GATGTGTCAA GTCTGCCGAT GCATGCTGGT AATGCTACTT ACGATACTGG AAGTTCGTTT 1740 GTTATCCCTC ATATTGATCA TATCCATGTC GTTCCGTATT CATGGTTGAC GCGCGATCAG 1800 ATTGCAACAA TCAAGTATGT GATGCAACAC CCCGAAGTTC GTCCAGATGT ATGGTCTAAG 1860 CCAGGCATG AAGAGTCAGG TTCGGTCATT CCAAATGTTA CGCCTCTTGA TAAACGTGCT 1920 GGTATGCCAA ATTGGCAAAT CATCCATTCT GCTGAAGAAG TTCAAAAAGC CCTAGCAGAA 1980 GGTCGTTTTG CAACACCAGA CGGCTATATT TTCGATCCAC GAGATGTTTT GGCCAAAGAA ACTTTTGTAT GGAAAGATGG CTCCTTTAGC ATCCCAAGAG CAGATGGCAG TTCATTGAGA ACCATTAATA AATCTGATCT ATCCCAAGCT GAGTGGCAAC AAGCTCAAGA GTTATTGGCA 2160 AAGAAAAACG CTGGTGATGC TACTGATACG GATAAACCCA AAGAAAAGCA ACAGGCAGAT 2220 AAGAGCAATG AAAACCAACA GCCAGTGAA GCCAGTAAAG AAGAAGAAAA AGAATCAGAT GACTTTATAG ACAGTTTACC AGACTATGGT CTAGATAGAG CAACCCTAGA AGATCATATC 2340 AATCAATTAG CACAAAAAGC TAATATCGAT CCTAAGTATC TCATTTTCCA ACCAGAAGGT 2400 GTCCAATTTT ATAATAAAAA TGGTGAATTA GTAACTTATG ATATCAAGAC GCTTCAACAA 2460 ATAAACCCTT AA (SEQ ID NO : 82) 2472

FIGURE 47

	VAAILLATHI		GLATKDNQIA	YIDDSKGKAK	50
APKTNKTMDQ	ISAEEGISAE	QIVVKITDQG	YVTSHGDHYH	FYNGKVPYDA	100
IISEELLMTD	~		KVNGNYYVYL	KPGSKRKNIR	150
	KGTKEAKEKG		EVAAVNEAKR	QGRYTTDDGY	200
					250
	APGRRKAPIP				300
	TFKELLDQLH				350
VPHGDHYHII	PRSQLSPLEM	ELADRYLAGQ	TEDNDSGSDH	SKPSDKEVTH	400
	GKGLDGKPYD				450
	LDEVANWVKA				500
VTKDGKVGYI	MPKDGKDYFY	ARDQLDLTQI	AFAEQELMLK	DKNHYRYDIV	550
DTGIEPRLAV	DVSSLPMHAG	NATYDTGSSF	VIPHIDHIHV	VPYSWLTRDQ	600
IATIKYVMQH	PEVRPDVWSK	PGHEESGSVI	PNVTPLDKRA	GMPNWQIIHS	650
AEEVQKALAE	GRFATPDGYI	FDPRDVLAKE	TFVWKDGSFS	IPRADGSSLR	700
TINKSDLSQA	EWQQAQELLA	KKNAGDATDT	DKPKEKQQAD	KSNENQQPSE	750
ASKEEEKESD	DFIDSLPDYG	LDRATLEDHI	NQLAQKANID	PKYLIFQPEG	800
VQFYNKNGEL	VTYDIKTLQQ	INPP (SEQ	ID NO : 83	3)	824

FIGURE 48